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(54) Title: ISOLATED HUMAN TRANSPORTER PROTEINS, NUCLEIC ACID MOLECULES ENCODING HUMAN TRANSPORTER PROTEINS, AND USES THEREOF

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(57) Abstract: The present invention provides amino acid sequences of peptides that are encoded by genes within the human genome, the transporter peptides of the present invention. The present invention specifically provides isolated peptide and nucleic acid molecules, methods of identifying orthologs and paralogs of the transporter peptides, and methods of identifying modulators of the transporter peptides.

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# ISOLATED HUMAN TRANSPORTER PROTEINS, NUCLEIC ACID MOLECULES ENCODING HUMAN TRANSPORTER PROTEINS, AND USES THEREOF

#### RELATED APPLICATIONS

The present application claims priority to provisional application U.S. Serial No. 60/240,836, filed October 17, 2000 (Atty. Docket CL000891-PROV) and 09/804,474, filed March 13, 2001(Atty. Docket CL000891).

#### FIELD OF THE INVENTION

The present invention is in the field of transporter proteins that are related to the sodium/calcium exchanger subfamily, recombinant DNA molecules, and protein production. The present invention specifically provides novel peptides and proteins that effect ligand transport and nucleic acid molecules encoding such peptide and protein molecules, all of which are useful in the development of human therapeutics and diagnostic compositions and methods.

#### **BACKGROUND OF THE INVENTION**

#### **Transporters**

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Transporter proteins regulate many different functions of a cell, including cell proliferation, differentiation, and signaling processes, by regulating the flow of molecules such as ions and macromolecules, into and out of cells. Transporters are found in the plasma membranes of virtually every cell in eukaryotic organisms. Transporters mediate a variety of cellular functions including regulation of membrane potentials and absorption and secretion of molecules and ion across cell membranes. When present in intracellular membranes of the Golgi apparatus and endocytic vesicles, transporters, such as chloride channels, also regulate organelle pH. For a review, see Greger, R. (1988) Annu. Rev. Physiol. 50:111-122.

Transporters are generally classified by structure and the type of mode of action. In addition, transporters are sometimes classified by the molecule type that is transported, for example, sugar transporters, chlorine channels, potassium channels, etc. There may be many classes of channels for transporting a single type of molecule (a detailed review of channel types can be found at Alexander, S.P.H. and J.A. Peters: Receptor and transporter nomenclature

supplement. Trends Pharmacol. Sci., Elsevier, pp. 65-68 (1997) and <a href="http://www-biology.ucsd.edu/~msaier/transport/titlepage2.html">http://www-biology.ucsd.edu/~msaier/transport/titlepage2.html</a>.

The following general classification scheme is known in the art and is followed in the present discoveries.

Channel-type transporters. Transmembrane channel proteins of this class are ubiquitously found in the membranes of all types of organisms from bacteria to higher eukaryotes. Transport systems of this type catalyze facilitated diffusion (by an energy-independent process) by passage through a transmembrane aqueous pore or channel without evidence for a carrier-mediated mechanism. These channel proteins usually consist largely of a-helical spanners, although b-strands may also be present and may even comprise the channel. However, outer membrane porin-type channel proteins are excluded from this class and are instead included in class 9.

Carrier-type transporters. Transport systems are included in this class if they utilize a carrier-mediated process to catalyze uniport (a single species is transported by facilitated diffusion), antiport (two or more species are transported in opposite directions in a tightly coupled process, not coupled to a direct form of energy other than chemiosmotic energy) and/or symport (two or more species are transported together in the same direction in a tightly coupled process, not coupled to a direct form of energy other than chemiosmotic energy).

Pyrophosphate bond hydrolysis-driven active transporters. Transport systems are included in this class if they hydrolyze pyrophosphate or the terminal pyrophosphate bond in ATP or another nucleoside triphosphate to drive the active uptake and/or extrusion of a solute or solutes. The transport protein may or may not be transiently phosphorylated, but the substrate is not phosphorylated.

PEP-dependent, phosphoryl transfer-driven group translocators. Transport systems of the bacterial phosphoenolpyruvate:sugar phosphotransferase system are included in this class. The product of the reaction, derived from extracellular sugar, is a cytoplasmic sugar-phosphate.

Decarboxylation-driven active transporters. Transport systems that drive solute (e.g., ion) uptake or extrusion by decarboxylation of a cytoplasmic substrate are included in this class.

Oxidoreduction-driven active transporters. Transport systems that drive transport of a solute (e.g., an ion) energized by the flow of electrons from a reduced substrate to an oxidized substrate are included in this class.

Light-driven active transporters. Transport systems that utilize light energy to drive transport of a solute (e.g., an ion) are included in this class.

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Mechanically-driven active transporters. Transport systems are included in this class if they drive movement of a cell or organelle by allowing the flow of ions (or other solutes) through the membrane down their electrochemical gradients.

Outer-membrane porins (of b-structure). These proteins form transmembrane pores or channels that usually allow the energy independent passage of solutes across a membrane. The transmembrane portions of these proteins consist exclusively of b-strands that form a b-barrel. These porin-type proteins are found in the outer membranes of Gram-negative bacteria, mitochondria and eukaryotic plastids.

Methyltransferase-driven active transporters. A single characterized protein currently falls into this category, the Na+-transporting methyltetrahydromethanopterin:coenzyme M methyltransferase.

Non-ribosome-synthesized channel-forming peptides or peptide-like molecules. These molecules, usually chains of L- and D-amino acids as well as other small molecular building blocks such as lactate, form oligomeric transmembrane ion channels. Voltage may induce channel formation by promoting assembly of the transmembrane channel. These peptides are often made by bacteria and fungi as agents of biological warfare.

Non-Proteinaceous Transport Complexes. Ion conducting substances in biological membranes that do not consist of or are not derived from proteins or peptides fall into this category.

Functionally characterized transporters for which sequence data are lacking. Transporters of particular physiological significance will be included in this category even though a family assignment cannot be made.

Putative transporters in which no family member is an established transporter. Putative transport protein families are grouped under this number and will either be classified elsewhere when the transport function of a member becomes established, or will be eliminated from the TC classification system if the proposed transport function is disproven. These families include a member or members for which a transport function has been suggested, but evidence for such a function is not yet compelling.

Auxiliary transport proteins. Proteins that in some way facilitate transport across one or more biological membranes but do not themselves participate directly in transport are included in this class. These proteins always function in conjunction with one or more transport proteins. They may provide a function connected with energy coupling to transport, play a structural role in complex formation or serve a regulatory function.

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Transporters of unknown classification. Transport protein families of unknown classification are grouped under this number and will be classified elsewhere when the transport process and energy coupling mechanism are characterized. These families include at least one member for which a transport function has been established, but either the mode of transport or the energy coupling mechanism is not known.

#### . Ion channels

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An important type of transporter is the ion channel. Ion channels regulate many different cell proliferation, differentiation, and signaling processes by regulating the flow of ions into and out of cells. Ion channels are found in the plasma membranes of virtually every cell in eukaryotic organisms. Ion channels mediate a variety of cellular functions including regulation of membrane potentials and absorption and secretion of ion across epithelial membranes. When present in intracellular membranes of the Golgi apparatus and endocytic vesicles, ion channels, such as chloride channels, also regulate organelle pH. For a review, see Greger, R. (1988) Annu. Rev. Physiol. 50:111-122.

Ion channels are generally classified by structure and the type of mode of action. For example, extracellular ligand gated channels (ELGs) are comprised of five polypeptide subunits, with each subunit having 4 membrane spanning domains, and are activated by the binding of an extracellular ligand to the channel. In addition, channels are sometimes classified by the ion type that is transported, for example, chlorine channels, potassium channels, etc. There may be many classes of channels for transporting a single type of ion (a detailed review of channel types can be found at Alexander, S.P.H. and J.A. Peters (1997). Receptor and ion channel nomenclature supplement. Trends Pharmacol. Sci., Elsevier, pp. 65-68 and http://www-biology.ucsd.edu/~msaier/transport/toc.html.

There are many types of ion channels based on structure. For example, many ion channels fall within one of the following groups: extracellular ligand-gated channels (ELG), intracellular ligand-gated channels (ILG), inward rectifying channels (INR), intercellular (gap junction) channels, and voltage gated channels (VIC). There are additionally recognized other channel families based on ion-type transported, cellular location and drug sensitivity. Detailed information on each of these, their activity, ligand type, ion type, disease association, drugability, and other information pertinent to the present invention, is well known in the art.

Extracellular ligand-gated channels, ELGs, are generally comprised of five polypeptide subunits, Unwin, N. (1993), Cell 72: 31-41; Unwin, N. (1995), Nature 373: 37-43; Hucho, F., et

al., (1996) J. Neurochem. 66: 1781-1792; Hucho, F., et al., (1996) Eur. J. Biochem. 239: 539-557; Alexander, S.P.H. and J.A. Peters (1997), Trends Pharmacol. Sci., Elsevier, pp. 4-6; 36-40; 42-44; and Xue, H. (1998) J. Mol. Evol. 47: 323-333. Each subunit has 4 membrane spanning regions: this serves as a means of identifying other members of the ELG family of proteins. ELG bind a ligand and in response modulate the flow of ions. Examples of ELG include most members of the neurotransmitter-receptor family of proteins, e.g., GABAI receptors. Other members of this family of ion channels include glycine receptors, ryandyne receptors, and ligand gated calcium channels.

#### Sodium/Calcium Exchangers

The protein provided by the present invention is a novel sodium/calcium exchanger. Sodium/calcium exchangers (NCX) rapidly import calcium during excitation impulse. Intracellular calcium concentrations vary greatly during the excitation/relaxation cycle. In contrast, extracellular calcium concentrations are maintained at relatively steady levels, despite wide variations in the amounts of calcium supplied with food.

There are at least three known mammalian NCX genes and a number of alternatively spliced isoforms. NCX sequences are highly conserved. NCX proteins contain 9 transmembrane domains and are regulated by calcium and sodium ions and, to some extent, by phosphorylation.

NCX proteins initiate cardiac myocyte contractions; this effect has been confirmed by *in vitro* experiments. Together with calsequestrin, a calcium binding protein, NCX proteins maintain calcium homeostasis in the heart muscle. This regulatory mechanism depends on the gene dosage, as evident from experiments with transgenic animals. Variations in expression levels of these proteins may be associated with some forms of heart disease.

Calcium transporters can mediate divalent ion toxicity. Barium and strontium can be carried by these channels into the cell, albeit at slower rates than calcium, which is the natural substrate. A panel of bivalent cations, such as copper, lead, cadmium, cobalt and nickel, inhibit calcium flow, but do not penetrate the cell membrane. Bivalent and trivalent iron, manganese, and zinc show no effect.

The sequence of the sodium/calcium exchanger provided by the present invention may be used to screen human populations for mutations associated with neurological conditions and heart disease. Furthermore, drugs can be designed that target this and other transporters.

For a further review of sodium/calcium exchangers, see: Linck et al., J Pharmacol Exp Ther 2000 Aug;294(2):648-57; Shen et al., J Pharmacol Exp Ther 2000 Aug;294(2):562-70;

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Philipson et al., Annu Rev Physiol 2000;62:111-33; Zhang et al., Br J Pharmacol 2000 Jun;130(3):485-8; and Vercesi et al., FEBS Lett 2000 May 12;473(2):203-6.

## The Voltage-gated Ion Channel (VIC) Superfamily

Proteins of the VIC family are ion-selective channel proteins found in a wide range of bacteria, archaea and eukaryotes Hille, B. (1992), Chapter 9: Structure of channel proteins; Chapter 20: Evolution and diversity. In: Ionic Channels of Excitable Membranes, 2nd Ed., Sinaur Assoc. Inc., Pubs., Sunderland, Massachusetts; Sigworth, F.J. (1993), Quart. Rev. Biophys. 27: 1-40; Salkoff, L. and T. Jegla (1995), Neuron 15: 489-492; Alexander, S.P.H. et al., (1997), Trends Pharmacol. Sci., Elsevier, pp. 76-84; Jan, L.Y. et al., (1997), Annu. Rev. Neurosci. 20: 91-123; Doyle, D.A, et al., (1998) Science 280: 69-77; Terlau, H. and W. Stühmer (1998), Naturwissenschaften 85: 437-444. They are often homo- or heterooligomeric structures with several dissimilar subunits (e.g., a1-a2-d-b Ca2+ channels, ab1b2 Na+ channels or (a)4-b K+ channels), but the channel and the primary receptor is usually associated with the a (or a1) subunit. Functionally characterized members are specific for K<sup>+</sup>, Na<sup>+</sup> or Ca<sup>2+</sup>. The K<sup>+</sup> channels usually consist of homotetrameric structures with each a-subunit possessing six transmembrane spanners (TMSs). The al and a subunits of the Ca2+ and Na+ channels, respectively, are about four times as large and possess 4 units, each with 6 TMSs separated by a hydrophilic loop, for a total of 24 TMSs. These large channel proteins form heterotetra-unit structures equivalent to the homotetrameric structures of most K<sup>+</sup> channels. All four units of the Ca<sup>2+</sup> and Na<sup>+</sup> channels are homologous to the single unit in the homotetrameric K<sup>+</sup> channels. Ion flux via the eukaryotic channels is generally controlled by the transmembrane electrical potential (hence the designation, voltage-sensitive) although some are controlled by ligand or receptor binding.

Several putative  $K^+$ -selective channel proteins of the VIC family have been identified in prokaryotes. The structure of one of them, the KcsA  $K^+$  channel of *Streptomyces lividans*, has been solved to 3.2 Å resolution. The protein possesses four identical subunits, each with two transmembrane helices, arranged in the shape of an inverted teepee or cone. The cone cradles the "selectivity filter" P domain in its outer end. The narrow selectivity filter is only 12 Å long, whereas the remainder of the channel is wider and lined with hydrophobic residues. A large water-filled cavity and helix dipoles stabilize  $K^+$  in the pore. The selectivity filter has two bound  $K^+$  ions about 7.5 Å apart from each other. Ion conduction is proposed to result from a balance of electrostatic attractive and repulsive forces.

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In eukaryotes, each VIC family channel type has several subtypes based on pharmacological and electrophysiological data. Thus, there are five types of Ca<sup>2+</sup> channels (L, N, P. O and T). There are at least ten types of K<sup>+</sup> channels, each responding in different ways to different stimuli: voltage-sensitive [Ka, Kv, Kvr, Kvs and Ksr], Ca2+-sensitive [BKCa, IKCa and SK<sub>Ca</sub>] and receptor-coupled [K<sub>M</sub> and K<sub>ACh</sub>]. There are at least six types of Na<sup>+</sup> channels (I, II, III, μ1, H1 and PN3). Tetrameric channels from both prokaryotic and eukaryotic organisms are known in which each a-subunit possesses 2 TMSs rather than 6, and these two TMSs are homologous to TMSs 5 and 6 of the six TMS unit found in the voltage-sensitive channel proteins. KcsA of S. lividans is an example of such a 2 TMS channel protein. These channels may include the K<sub>Na</sub> (Na<sup>+</sup>-activated) and K<sub>Vol</sub> (cell volume-sensitive) K<sup>+</sup> channels, as well as distantly related channels such as the Tok1 K<sup>+</sup> channel of yeast, the TWIK-1 inward rectifier K<sup>+</sup> channel of the mouse and the TREK-1 K<sup>+</sup> channel of the mouse. Because of insufficient sequence similarity with proteins of the VIC family, inward rectifier K<sup>+</sup> IRK channels (ATPregulated; G-protein-activated) which possess a P domain and two flanking TMSs are placed in a distinct family. However, substantial sequence similarity in the P region suggests that they are homologous. The b, g and d subunits of VIC family members, when present, frequently play regulatory roles in channel activation/deactivation.

## The Epithelial Na<sup>+</sup> Channel (ENaC) Family

The ENaC family consists of over twenty-four sequenced proteins (Canessa, C.M., et al., (1994), Nature 367: 463-467, Le, T. and M.H. Saier, Jr. (1996), Mol. Membr. Biol. 13: 149-157; Garty, H. and L.G. Palmer (1997), Physiol. Rev. 77: 359-396; Waldmann, R., et al., (1997), Nature 386: 173-177; Darboux, I., et al., (1998), J. Biol. Chem. 273: 9424-9429; Firsov, D., et al., (1998), EMBO J. 17: 344-352; Horisberger, J.-D. (1998). Curr. Opin. Struc. Biol. 10: 443-449). All are from animals with no recognizable homologues in other eukaryotes or bacteria.

The vertebrate ENaC proteins from epithelial cells cluster tightly together on the phylogenetic tree: voltage-insensitive ENaC homologues are also found in the brain. Eleven sequenced C. elegans proteins, including the degenerins, are distantly related to the vertebrate proteins as well as to each other. At least some of these proteins form part of a mechano-transducing complex for touch sensitivity. The homologous Helix aspersa (FMRF-amide)-activated Na<sup>+</sup> channel is the first peptide neurotransmitter-gated ionotropic receptor to be sequenced.

Protein members of this family all exhibit the same apparent topology, each with N- and C-termini on the inside of the cell, two amphipathic transmembrane spanning segments, and a

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large extracellular loop. The extracellular domains contain numerous highly conserved cysteine residues. They are proposed to serve a receptor function.

Mammalian ENaC is important for the maintenance of Na<sup>+</sup> balance and the regulation of blood pressure. Three homologous ENaC subunits, alpha, beta, and gamma, have been shown to assemble to form the highly Na <sup>+</sup>-selective channel. The stoichiometry of the three subunits is alpha<sub>2</sub>, beta1, gamma1 in a heterotetrameric architecture.

## The Glutamate-gated Ion Channel (GIC) Family of Neurotransmitter Receptors

Members of the GIC family are heteropentameric complexes in which each of the 5 subunits is of 800-1000 amino acyl residues in length (Nakanishi, N., et al, (1990), Neuron 5: 569-581; Unwin, N. (1993), Cell 72: 31-41; Alexander, S.P.H. and J.A. Peters (1997) Trends Pharmacol. Sci., Elsevier, pp. 36-40). These subunits may span the membrane three or five times as putative a-helices with the N-termini (the glutamate-binding domains) localized extracellularly and the C-termini localized cytoplasmically. They may be distantly related to the ligand-gated ion channels, and if so, they may possess substantial b-structure in their transmembrane regions. However, homology between these two families cannot be established on the basis of sequence comparisons alone. The subunits fall into six subfamilies: a, b, g, d, e and z.

The GIC channels are divided into three types: (1) a-amino-3-hydroxy-5-methyl-4-isoxazole propionate (AMPA)-, (2) kainate- and (3) N-methyl-D-aspartate (NMDA)-selective glutamate receptors. Subunits of the AMPA and kainate classes exhibit 35-40% identity with each other while subunits of the NMDA receptors exhibit 22-24% identity with the former subunits. They possess large N-terminal, extracellular glutamate-binding domains that are homologous to the periplasmic glutamine and glutamate receptors of ABC-type uptake permeases of Gram-negative bacteria. All known members of the GIC family are from animals. The different channel (receptor) types exhibit distinct ion selectivities and conductance properties. The NMDA-selective large conductance channels are highly permeable to monovalent cations and Ca<sup>2+</sup>. The AMPA- and kainate-selective ion channels are permeable primarily to monovalent cations with only low permeability to Ca<sup>2+</sup>.

#### The Chloride Channel (CIC) Family

The ClC family is a large family consisting of dozens of sequenced proteins derived from Gram-negative and Gram-positive bacteria, cyanobacteria, archaea, yeast, plants and animals (Steinmeyer, K., et al., (1991), Nature 354: 301-304; Uchida, S., et al., (1993), J. Biol. Chem.

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268: 3821-3824; Huang, M.-E., et al., (1994), J. Mol. Biol. 242: 595-598; Kawasaki, M., et al, (1994), Neuron 12: 597-604; Fisher, W.E., et al., (1995), Genomics. 29:598-606; and Foskett, J.K. (1998), Annu. Rev. Physiol. 60: 689-717). These proteins are essentially ubiquitous, although they are not encoded within genomes of Haemophilus influenzae, Mycoplasma genitalium, and Mycoplasma pneumoniae. Sequenced proteins vary in size from 395 amino acyl residues (M. jannaschii) to 988 residues (man). Several organisms contain multiple ClC family paralogues. For example, Synechocystis has two paralogues, one of 451 residues in length and the other of 899 residues. Arabidopsis thaliana has at least four sequenced paralogues, (775-792 residues), humans also have at least five paralogues (820-988 residues), and C. elegans also has at least five (810-950 residues). There are nine known members in mammals, and mutations in three of the corresponding genes cause human diseases. E. coli, Methanococcus jannaschii and Saccharomyces cerevisiae only have one ClC family member each. With the exception of the larger Synechocystis paralogue, all bacterial proteins are small (395-492 residues) while all eukaryotic proteins are larger (687-988 residues). These proteins exhibit 10-12 putative transmembrane a-helical spanners (TMSs) and appear to be present in the membrane as homodimers. While one member of the family, Torpedo ClC-O, has been reported to have two channels, one per subunit, others are believed to have just one.

All functionally characterized members of the ClC family transport chloride, some in a voltage-regulated process. These channels serve a variety of physiological functions (cell volume regulation; membrane potential stabilization; signal transduction; transepithelial transport, etc.). Different homologues in humans exhibit differing anion selectivities, i.e., ClC4 and ClC5 share a  $NO_3^- > Cl^- > Br^- > I^-$  conductance sequence, while ClC3 has an  $I^- > Cl^-$  selectivity. The ClC4 and ClC5 channels and others exhibit outward rectifying currents with currents only at voltages more positive than +20mV.

#### Animal Inward Rectifier K<sup>+</sup> Channel (IRK-C) Family

IRK channels possess the "minimal channel-forming structure" with only a P domain, characteristic of the channel proteins of the VIC family, and two flanking transmembrane spanners (Shuck, M.E., et al., (1994), J. Biol. Chem. 269: 24261-24270; Ashen, M.D., et al., (1995), Am. J. Physiol. 268: H506-H511; Salkoff, L. and T. Jegla (1995), Neuron 15: 489-492; Aguilar-Bryan, L., et al., (1998), Physiol. Rev. 78: 227-245; Ruknudin, A., et al., (1998), J. Biol. Chem. 273: 14165-14171). They may exist in the membrane as homo- or heterooligomers. They have a greater tendency to let K<sup>+</sup> flow into the cell than out. Voltage-dependence may be regulated by external K<sup>+</sup>, by internal Mg<sup>2+</sup>, by internal ATP and/or by G-proteins. The P domains

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of IRK channels exhibit limited sequence similarity to those of the VIC family, but this sequence similarity is insufficient to establish homology. Inward rectifiers play a role in setting cellular membrane potentials, and the closing of these channels upon depolarization permits the occurrence of long duration action potentials with a plateau phase. Inward rectifiers lack the intrinsic voltage sensing helices found in VIC family channels. In a few cases, those of Kirl.la and Kir6.2, for example, direct interaction with a member of the ABC superfamily has been proposed to confer unique functional and regulatory properties to the heteromeric complex, including sensitivity to ATP. The SUR1 sulfonylurea receptor (spQ09428) is the ABC protein that regulates the Kir6.2 channel in response to ATP, and CFTR may regulate Kirl.la. Mutations in SUR1 are the cause of familial persistent hyperinsulinemic hypoglycemia in infancy (PHHI), an autosomal recessive disorder characterized by unregulated insulin secretion in the pancreas.

#### ATP-gated Cation Channel (ACC) Family

Members of the ACC family (also called P2X receptors) respond to ATP, a functional neurotransmitter released by exocytosis from many types of neurons (North, R.A. (1996), Curr. Opin. Cell Biol. 8: 474-483; Soto, F., M. Garcia-Guzman and W. Stühmer (1997), J. Membr. Biol. 160: 91-100). They have been placed into seven groups (P2X<sub>1</sub> - P2X<sub>7</sub>) based on their pharmacological properties. These channels, which function at neuron-neuron and neuron-smooth muscle junctions, may play roles in the control of blood pressure and pain sensation. They may also function in lymphocyte and platelet physiology. They are found only in animals.

The proteins of the ACC family are quite similar in sequence (>35% identity), but they possess 380-1000 amino acyl residues per subunit with variability in length localized primarily to the C-terminal domains. They possess two transmembrane spanners, one about 30-50 residues from their N-termini, the other near residues 320-340. The extracellular receptor domains between these two spanners (of about 270 residues) are well conserved with numerous conserved glycyl and cysteyl residues. The hydrophilic C-termini vary in length from 25 to 240 residues. They resemble the topologically similar epithelial Na<sup>+</sup> channel (ENaC) proteins in possessing (a) N- and C-termini localized intracellularly, (b) two putative transmembrane spanners, (c) a large extracellular loop domain, and (d) many conserved extracellular cysteyl residues. ACC family members are, however, not demonstrably homologous with them. ACC channels are probably hetero- or homomultimers and transport small monovalent cations (Me<sup>+</sup>). Some also transport Ca<sup>2+</sup>; a few also transport small metabolites.

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The Ryanodine-Inositol 1,4,5-triphosphate Receptor Ca<sup>2+</sup> Channel (RIR-CaC) Family

Ryanodine (Ry)-sensitive and inositol 1,4,5-triphosphate (IP3)-sensitive Ca<sup>2+</sup>-release channels function in the release of Ca<sup>2+</sup> from intracellular storage sites in animal cells and thereby regulate various Ca<sup>2+</sup> -dependent physiological processes (Hasan, G. et al., (1992) Development 116: 967-975; Michikawa, T., et al., (1994), J. Biol. Chem. 269: 9184-9189; Tunwell, R.E.A., (1996), Biochem. J. 318: 477-487; Lee, A.G. (1996) *Biomembranes*, Vol. 6, Transmembrane Receptors and Channels (A.G. Lee, ed.), JAI Press, Denver, CO., pp 291-326; Mikoshiba, K., et al., (1996) J. Biochem. Biomem. 6: 273-289). Ry receptors occur primarily in muscle cell sarcoplasmic reticular (SR) membranes, and IP3 receptors occur primarily in brain cell endoplasmic reticular (FR) membranes where they effect release of Ca<sup>2+</sup> into the cytoplasm upon activation (opening) of the channel.

The Ry receptors are activated as a result of the activity of dihydropyridine-sensitive Ca<sup>2+</sup> channels. The latter are members of the voltage-sensitive ion channel (VIC) family.

Dihydropyridine-sensitive channels are present in the T-tubular systems of muscle tissues.

Ry receptors are homotetrameric complexes with each subunit exhibiting a molecular size of over 500,000 daltons (about 5,000 amino acyl residues). They possess C-terminal domains with six putative transmembrane a -helical spanners (TMSs). Putative pore-forming sequences occur between the fifth and sixth TMSs as suggested for members of the VIC family. The large N-terminal hydrophilic domains and the small C-terminal hydrophilic domains are localized to the cytoplasm. Low resolution 3-dimensional structural data are available. Mammals possess at least three isoforms that probably arose by gene duplication and divergence before divergence of the mammalian species. Homologues are present in humans and Caenorabditis elegans.

IP<sub>3</sub> receptors resemble Ry receptors in many respects. (1) They are homotetrameric complexes with each subunit exhibiting a molecular size of over 300,000 daltons (about 2,700 amino acyl residues). (2) They possess C-terminal channel domains that are homologous to those of the Ry receptors. (3) The channel domains possess six putative TMSs and a putative channel lining region between TMSs 5 and 6. (4) Both the large N-terminal domains and the smaller C-terminal tails face the cytoplasm. (5) They possess covalently linked carbohydrate on extracytoplasmic loops of the channel domains. (6) They have three currently recognized isoforms (types 1, 2, and 3) in mammals which are subject to differential regulation and have different tissue distributions.

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IP<sub>3</sub> receptors possess three domains: N-terminal IP<sub>3</sub>-binding domains, central coupling or regulatory domains and C-terminal channel domains. Channels are activated by IP<sub>3</sub> binding, and like the Ry receptors, the activities of the IP<sub>3</sub> receptor channels are regulated by phosphorylation of the regulatory domains, catalyzed by various protein kinases. They predominate in the endoplasmic reticular membranes of various cell types in the brain but have also been found in the plasma membranes of some nerve cells derived from a variety of tissues.

The channel domains of the Ry and IP<sub>3</sub> receptors comprise a coherent family that in spite of apparent structural similarities, do not show appreciable sequence similarity of the proteins of the VIC family. The Ry receptors and the IP<sub>3</sub> receptors cluster separately on the RIR-CaC family tree. They both have homologues in *Drosophila*. Based on the phylogenetic tree for the family, the family probably evolved in the following sequence: (1) A gene duplication event occurred that gave rise to Ry and IP<sub>3</sub> receptors in invertebrates. (2) Vertebrates evolved from invertebrates. (3) The three isoforms of each receptor arose as a result of two distinct gene duplication events. (4) These isoforms were transmitted to mammals before divergence of the mammalian species.

## The Organellar Chloride Channel (O-ClC) Family

Proteins of the O-ClC family are voltage-sensitive chloride channels found in intracellular membranes but not the plasma membranes of animal cells (Landry, D, et al., (1993), J. Biol. Chem. 268: 14948-14955; Valenzuela, Set al., (1997), J. Biol. Chem. 272: 12575-12582; and Duncan, R.R., et al., (1997), J. Biol. Chem. 272: 23880-23886).

They are found in human nuclear membranes, and the bovine protein targets to the microsomes, but not the plasma membrane, when expressed in *Xenopus laevis* oocytes. These proteins are thought to function in the regulation of the membrane potential and in transepithelial ion absorption and secretion in the kidney. They possess two putative transmembrane a-helical spanners (TMSs) with cytoplasmic N- and C-termini and a large luminal loop that may be glycosylated. The bovine protein is 437 amino acyl residues in length and has the two putative TMSs at positions 223-239 and 367-385. The human nuclear protein is much smaller (241 residues). A *C. elegans* homologue is 260 residues long.

Transporter proteins, particularly members of the sodium/calcium exchanger subfamily, are a major target for drug action and development. Accordingly, it is valuable to the field of pharmaceutical development to identify and characterize previously unknown transport proteins. The present invention advances the state of the art by providing previously unidentified human transport proteins.

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#### SUMMARY OF THE INVENTION

The present invention is based in part on the identification of amino acid sequences of human transporter peptides and proteins that are related to the sodium/calcium exchanger subfamily, as well as allelic variants and other mammalian orthologs thereof. These unique peptide sequences, and nucleic acid sequences that encode these peptides, can be used as models for the development of human therapeutic targets, aid in the identification of therapeutic proteins, and serve as targets for the development of human therapeutic agents that modulate transporter activity in cells and tissues that express the transporter. Experimental data as provided in Figure 1 indicates expression in humans in brain, heart, kidney, lung, spleen, testis, leukocyte and fetal brain.

#### DESCRIPTION OF THE FIGURE SHEETS

FIGURE 1 provides the nucleotide sequence of a cDNA molecule or transcript sequence that encodes the transporter protein of the present invention (SEQ ID NO:1). In addition structure and functional information is provided, such as ATG start, stop and tissue distribution, where available, that allows one to readily determine specific uses of inventions based on this molecular sequence. Experimental data as provided in Figure 1 indicates expression in humans in brain, heart, kidney, lung, spleen, testis, leukocyte and fetal brain.

FIGURE 2 provides the predicted amino acid sequence of the transporter of the present invention. (SEQ ID NO:2) In addition structure and functional information such as protein family, function, and modification sites is provided where available, allowing one to readily determine specific uses of inventions based on this molecular sequence.

FIGURE 3 provides genomic sequences that span the gene encoding the transporter protein of the present invention (SEQ ID NO: 3). In addition structure and functional information, such as intron/exon structure, promoter location, etc., is provided where available, allowing one to readily determine specific uses of inventions based on this molecular sequence. 140 SNPs, including 6 indels, have been identified in the gene encoding the transporter protein provided by the present invention and are given in Figure 3.

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## DETAILED DESCRIPTION OF THE INVENTION

#### General Description

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The present invention is based on the sequencing of the human genome. During the sequencing and assembly of the human genome, analysis of the sequence information revealed previously unidentified fragments of the human genome that encode peptides that share structural and/or sequence homology to protein/peptide/domains identified and characterized within the art as being a transporter protein or part of a transporter protein and are related to the sodium/calcium exchanger subfamily. Utilizing these sequences, additional genomic sequences were assembled and transcript and/or cDNA sequences were isolated and characterized. Based on this analysis, the present invention provides amino acid sequences of human transporter peptides and proteins that are related to the sodium/calcium exchanger subfamily, nucleic acid sequences in the form of transcript sequences, cDNA sequences and/or genomic sequences that encode these transporter peptides and proteins, nucleic acid variation (allelic information), tissue distribution of expression, and information about the closest art known protein/peptide/domain that has structural or sequence homology to the transporter of the present invention.

In addition to being previously unknown, the peptides that are provided in the present invention are selected based on their ability to be used for the development of commercially important products and services. Specifically, the present peptides are selected based on homology and/or structural relatedness to known transporter proteins of the sodium/calcium exchanger subfamily and the expression pattern observed. Experimental data as provided in Figure 1 indicates expression in humans in brain, heart, kidney, lung, spleen, testis, leukocyte and fetal brain. The art has clearly established the commercial importance of members of this family of proteins and proteins that have expression patterns similar to that of the present gene. Some of the more specific features of the peptides of the present invention, and the uses thereof, are described herein, particularly in the Background of the Invention and in the annotation provided in the Figures, and/or are known within the art for each of the known sodium/calcium exchanger family or subfamily of transporter proteins.

#### Specific Embodiments

#### Peptide Molecules

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The present invention provides nucleic acid sequences that encode protein molecules that have been identified as being members of the transporter family of proteins and are related to the sodiunt/calcium exchanger subfamily (protein sequences are provided in Figure 2, transcript/cDNA sequences are provided in Figures 1 and genomic sequences are provided in Figure 3). The peptide sequences provided in Figure 2, as well as the obvious variants described herein, particularly allelic variants as identified herein and using the information in Figure 3, will be referred herein as the transporter peptides of the present invention, transporter peptides, or peptides/proteins of the present invention.

The present invention provides isolated peptide and protein molecules that consist of, consist essentially of, or comprising the amino acid sequences of the transporter peptides disclosed in the Figure 2, (encoded by the nucleic acid molecule shown in Figure 1, transcript/cDNA or Figure 3, genomic sequence), as well as all obvious variants of these peptides that are within the art to make and use. Some of these variants are described in detail below.

As used herein, a peptide is said to be "isolated" or "purified" when it is substantially free of cellular material or free of chemical precursors or other chemicals. The peptides of the present invention can be purified to homogeneity or other degrees of purity. The level of purification will be based on the intended use. The critical feature is that the preparation allows for the desired function of the peptide, even if in the presence of considerable amounts of other components (the features of an isolated nucleic acid molecule is discussed below).

In some uses, "substantially free of cellular material" includes preparations of the peptide having less than about 30% (by dry weight) other proteins (i.e., contaminating protein), less than about 20% other proteins, less than about 10% other proteins, or less than about 5% other proteins. When the peptide is recombinantly produced, it can also be substantially free of culture medium, i.e., culture medium represents less than about 20% of the volume of the protein preparation.

The language "substantially free of chemical precursors or other chemicals" includes preparations of the peptide in which it is separated from chemical precursors or other chemicals that are involved in its synthesis. In one embodiment, the language "substantially free of chemical precursors or other chemicals" includes preparations of the transporter peptide having less than about 30% (by dry weight) chemical precursors or other chemicals, less than about 20% chemical

precursors or other chemicals, less than about 10% chemical precursors or other chemicals, or less than about 5% chemical precursors or other chemicals.

The isolated transporter peptide can be purified from cells that naturally express it, purified from cells that have been altered to express it (recombinant), or synthesized using known protein synthesis methods. Experimental data as provided in Figure 1 indicates expression in humans in brain, heart, kidney, lung, spleen, testis, leukocyte and fetal brain. For example, a nucleic acid molecule encoding the transporter peptide is cloned into an expression vector, the expression vector introduced into a host cell and the protein expressed in the host cell. The protein can then be isolated from the cells by an appropriate purification scheme using standard protein purification techniques. Many of these techniques are described in detail below.

Accordingly, the present invention provides proteins that consist of the amino acid sequences provided in Figure 2 (SEQ ID NO:2), for example, proteins encoded by the transcript/cDNA nucleic acid sequences shown in Figure 1 (SEQ ID NO:1) and the genomic sequences provided in Figure 3 (SEQ ID NO:3). The amino acid sequence of such a protein is provided in Figure 2. A protein consists of an amino acid sequence when the amino acid sequence is the final amino acid sequence of the protein.

The present invention further provides proteins that consist essentially of the amino acid sequences provided in Figure 2 (SEQ ID NO:2), for example, proteins encoded by the transcript/cDNA nucleic acid sequences shown in Figure 1 (SEQ ID NO:1) and the genomic sequences provided in Figure 3 (SEQ ID NO:3). A protein consists essentially of an amino acid sequence when such an amino acid sequence is present with only a few additional amino acid residues, for example from about 1 to about 100 or so additional residues, typically from 1 to about 20 additional residues in the final protein.

The present invention further provides proteins that comprise the amino acid sequences provided in Figure 2 (SEQ ID NO:2), for example, proteins encoded by the transcript/cDNA nucleic acid sequences shown in Figure 1 (SEQ ID NO:1) and the genomic sequences provided in Figure 3 (SEQ ID NO:3). A protein comprises an amino acid sequence when the amino acid sequence is at least part of the final amino acid sequence of the protein. In such a fashion, the protein can be only the peptide or have additional amino acid molecules, such as amino acid residues (contiguous encoded sequence) that are naturally associated with it or heterologous amino acid residues/peptide sequences. Such a protein can have a few additional amino acid residues or can comprise several hundred or more additional amino acids. The preferred classes of proteins that are comprised of the

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transporter peptides of the present invention are the naturally occurring mature proteins. A brief description of how various types of these proteins can be made/isolated is provided below.

The transporter peptides of the present invention can be attached to heterologous sequences to form chimeric or fusion proteins. Such chimeric and fusion proteins comprise a transporter peptide operatively linked to a heterologous protein having an amino acid sequence not substantially homologous to the transporter peptide. "Operatively linked" indicates that the transporter peptide and the heterologous protein are fused in-frame. The heterologous protein can be fused to the N-terminus or C-terminus of the transporter peptide.

In some uses, the fusion protein does not affect the activity of the transporter peptide per se. For example, the fusion protein can include, but is not limited to, enzymatic fusion proteins, for example beta-galactosidase fusions, yeast two-hybrid GAL fusions, poly-His fusions, MYC-tagged, HI-tagged and Ig fusions. Such fusion proteins, particularly poly-His fusions, can facilitate the purification of recombinant transporter peptide. In certain host cells (e.g., mammalian host cells), expression and/or secretion of a protein can be increased by using a heterologous signal sequence.

A chimeric or fusion protein can be produced by standard recombinant DNA techniques. For example, DNA fragments coding for the different protein sequences are ligated together inframe in accordance with conventional techniques. In another embodiment, the fusion gene can be synthesized by conventional techniques including automated DNA synthesizers. Alternatively, PCR amplification of gene fragments can be carried out using anchor primers which give rise to complementary overhangs between two consecutive gene fragments which can subsequently be annealed and re-amplified to generate a chimeric gene sequence (see Ausubel *et al.*, *Current Protocols in Molecular Biology*, 1992). Moreover, many expression vectors are commercially available that already encode a fusion moiety (e.g., a GST protein). A transporter peptide-encoding nucleic acid can be cloned into such an expression vector such that the fusion moiety is linked inframe to the transporter peptide.

As mentioned above, the present invention also provides and enables obvious variants of the amino acid sequence of the proteins of the present invention, such as naturally occurring mature forms of the peptide, allelic/sequence variants of the peptides, non-naturally occurring recombinantly derived variants of the peptides, and orthologs and paralogs of the peptides. Such variants can readily be generated using art-known techniques in the fields of recombinant nucleic acid technology and protein biochemistry. It is understood, however, that variants exclude any amino acid sequences disclosed prior to the invention.

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Such variants can readily be identified/made using molecular techniques and the sequence information disclosed herein. Further, such variants can readily be distinguished from other peptides based on sequence and/or structural homology to the transporter peptides of the present invention. The degree of homology/identity present will be based primarily on whether the peptide is a functional variant or non-functional variant, the amount of divergence present in the paralog family and the evolutionary distance between the orthologs.

To determine the percent identity of two amino acid sequences or two nucleic acid sequences, the sequences are aligned for optimal comparison purposes (e.g., gaps can be introduced in one or both of a first and a second amino acid or nucleic acid sequence for optimal alignment and non-homologous sequences can be disregarded for comparison purposes). In a preferred embodiment, at least 30%, 40%, 50%, 60%, 70%, 80%, or 90% or more of a reference sequence is aligned for comparison purposes. The amino acid residues or nucleotides at corresponding amino acid positions or nucleotide positions are then compared. When a position in the first sequence is occupied by the same amino acid residue or nucleotide as the corresponding position in the second sequence, then the molecules are identical at that position (as used herein amino acid or nucleic acid "identity" is equivalent to amino acid or nucleic acid "homology"). The percent identity between the two sequences is a function of the number of identical positions shared by the sequences, taking into account the number of gaps, and the length of each gap, which need to be introduced for optimal alignment of the two sequences.

The comparison of sequences and determination of percent identity and similarity between two sequences can be accomplished using a mathematical algorithm. (Computational Molecular Biology, Lesk, A.M., ed., Oxford University Press, New York, 1988; Biocomputing: Informatics and Genome Projects, Smith, D.W., ed., Academic Press, New York, 1993; Computer Analysis of Sequence Data, Part 1, Griffin, A.M., and Griffin, H.G., eds., Humana Press, New Jersey, 1994; Sequence Analysis in Molecular Biology, von Heinje, G., Academic Press, 1987; and Sequence Analysis Primer, Gribskov, M. and Devereux, J., eds., M Stockton Press, New York, 1991). In a preferred embodiment, the percent identity between two amino acid sequences is determined using the Needleman and Wunsch (J. Mol. Biol. (48):444-453 (1970)) algorithm which has been incorporated into the GAP program in the GCG software package (available at http://www.gcg.com), using either a Blossom 62 matrix or a PAM250 matrix, and a gap weight of 16, 14, 12, 10, 8, 6, or 4 and a length weight of 1, 2, 3, 4, 5, or 6. In yet another preferred embodiment, the percent identity between two nucleotide sequences is determined using the GAP program in the GCG software package (Devereux, J., et al., Nucleic Acids Res. 12(1):387

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(1984)) (available at http://www.gcg.com), using a NWSgapdna.CMP matrix and a gap weight of 40, 50, 60, 70, or 80 and a length weight of 1, 2, 3, 4, 5, or 6. In another embodiment, the percent identity between two amino acid or nucleotide sequences is determined using the algorithm of E. Myers and W. Miller (CABIOS, 4:11-17 (1989)) which has been incorporated into the ALIGN program (version 2.0), using a PAM120 weight residue table, a gap length penalty of 12 and a gap penalty of 4.

The nucleic acid and protein sequences of the present invention can further be used as a "query sequence" to perform a search against sequence databases to, for example, identify other family members or related sequences. Such searches can be performed using the NBLAST and XBLAST programs (version 2.0) of Altschul, et al. (J. Mol. Biol. 215:403-10 (1990)). BLAST nucleotide searches can be performed with the NBLAST program, score = 100, wordlength = 12 to obtain nucleotide sequences homologous to the nucleic acid molecules of the invention.

BLAST protein searches can be performed with the XBLAST program, score = 50, wordlength = 3 to obtain amino acid sequences homologous to the proteins of the invention. To obtain gapped alignments for comparison purposes, Gapped BLAST can be utilized as described in Altschul et al. (Nucleic Acids Res. 25(17):3389-3402 (1997)). When utilizing BLAST and gapped BLAST programs, the default parameters of the respective programs (e.g., XBLAST and NBLAST) can be used.

Full-length pre-processed forms, as well as mature processed forms, of proteins that comprise one of the peptides of the present invention can readily be identified as having complete sequence identity to one of the transporter peptides of the present invention as well as being encoded by the same genetic locus as the transporter peptide provided herein. As indicated by the data presented in Figure 3, the map position was determined to be on chromosome 14 by ePCR.

Allelic variants of a transporter peptide can readily be identified as being a human protein having a high degree (significant) of sequence homology/identity to at least a portion of the transporter peptide as well as being encoded by the same genetic locus as the transporter peptide provided herein. Genetic locus can readily be determined based on the genomic information provided in Figure 3, such as the genomic sequence mapped to the reference human. As indicated by the data presented in Figure 3, the map position was determined to be on chromosome 14 by ePCR. As used herein, two proteins (or a region of the proteins) have significant homology when the amino acid sequences are typically at least about 70-80%, 80-90%, and more typically at least about 90-95% or more homologous. A significantly homologous amino acid sequence, according to the present invention, will be encoded by a nucleic acid sequence that will hybridize

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to a transporter peptide encoding nucleic acid molecule under stringent conditions as more fully described below.

Figure 3 provides information on SNPs that have been identified in a gene encoding the transporter protein of the present invention. 140 SNP variants were found, including 6 indels (indicated by a "-") and 1 SNPs in exons. The others were found in in introns and regions 5' and 3' of the ORF. Such SNPs in introns and outside the ORF may affect control/regulatory elements.

Paralogs of a transporter peptide can readily be identified as having some degree of significant sequence homology/identity to at least a portion of the transporter peptide, as being encoded by a gene from humans, and as having similar activity or function. Two proteins will typically be considered paralogs when the amino acid sequences are typically at least about 60% or greater, and more typically at least about 70% or greater homology through a given region or domain. Such paralogs will be encoded by a nucleic acid sequence that will hybridize to a transporter peptide encoding nucleic acid molecule under moderate to stringent conditions as more fully described below.

Orthologs of a transporter peptide can readily be identified as having some degree of significant sequence homology/identity to at least a portion of the transporter peptide as well as being encoded by a gene from another organism. Preferred orthologs will be isolated from mammals, preferably primates, for the development of human therapeutic targets and agents. Such orthologs will be encoded by a nucleic acid sequence that will hybridize to a transporter peptide encoding nucleic acid molecule under moderate to stringent conditions, as more fully described below, depending on the degree of relatedness of the two organisms yielding the proteins.

Non-naturally occurring variants of the transporter peptides of the present invention can readily be generated using recombinant techniques. Such variants include, but are not limited to deletions, additions and substitutions in the amino acid sequence of the transporter peptide. For example, one class of substitutions are conserved amino acid substitution. Such substitutions are those that substitute a given amino acid in a transporter peptide by another amino acid of like characteristics. Typically seen as conservative substitutions are the replacements, one for another, among the aliphatic amino acids Ala, Val, Leu, and Ile; interchange of the hydroxyl residues Ser and Thr; exchange of the acidic residues Asp and Glu; substitution between the amide residues Asp and Gln; exchange of the basic residues Lys and Arg; and replacements among the aromatic residues Phe and Tyr. Guidance concerning which amino acid changes are likely to be phenotypically silent are found in Bowie *et al.*, *Science 247*:1306-1310 (1990).

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Variant transporter peptides can be fully functional or can lack function in one or more activities, e.g. ability to bind ligand, ability to transport ligand, ability to mediate signaling, etc. Fully functional variants typically contain only conservative variation or variation in non-critical residues or in non-critical regions. Figure 2 provides the result of protein analysis and can be used to identify critical domains/regions. Functional variants can also contain substitution of similar amino acids that result in no change or an insignificant change in function. Alternatively, such substitutions may positively or negatively affect function to some degree.

Non-functional variants typically contain one or more non-conservative amino acid substitutions, deletions, insertions, inversions, or truncation or a substitution, insertion, inversion, or deletion in a critical residue or critical region.

Amino acids that are essential for function can be identified by methods known in the art, such as site-directed mutagenesis or alanine-scanning mutagenesis (Cunningham et al., Science 244:1081-1085 (1989)), particularly using the results provided in Figure 2. The latter procedure introduces single alanine mutations at every residue in the molecule. The resulting mutant molecules are then tested for biological activity such as transporter activity or in assays such as an in vitro proliferative activity. Sites that are critical for binding partner/substrate binding can also be determined by structural analysis such as crystallization, nuclear magnetic resonance or photoaffinity labeling (Smith et al., J. Mol. Biol. 224:899-904 (1992); de Vos et al. Science 255:306-312 (1992)).

The present invention further provides fragments of the transporter peptides, in addition to proteins and peptides that comprise and consist of such fragments, particularly those comprising the residues identified in Figure 2. The fragments to which the invention pertains, however, are not to be construed as encompassing fragments that may be disclosed publicly prior to the present invention.

As used herein, a fragment comprises at least 8, 10, 12, 14, 16, or more contiguous amino acid residues from a transporter peptide. Such fragments can be chosen based on the ability to retain one or more of the biological activities of the transporter peptide or could be chosen for the ability to perform a function, e.g. bind a substrate or act as an immunogen. Particularly important fragments are biologically active fragments, peptides that are, for example, about 8 or more amino acids in length. Such fragments will typically comprise a domain or motif of the transporter peptide, e.g., active site, a transmembrane domain or a substrate-binding domain. Further, possible fragments include, but are not limited to, domain or motif containing fragments, soluble peptide fragments, and fragments containing immunogenic structures. Predicted domains and functional

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sites are readily identifiable by computer programs well known and readily available to those of skill in the art (e.g., PROSITE analysis). The results of one such analysis are provided in Figure 2.

Polypeptides often contain amino acids other than the 20 amino acids commonly referred to as the 20 naturally occurring amino acids. Further, many amino acids, including the terminal amino acids, may be modified by natural processes, such as processing and other post-translational modifications, or by chemical modification techniques well known in the art. Common modifications that occur naturally in transporter peptides are described in basic texts, detailed monographs, and the research literature, and they are well known to those of skill in the art (some of these features are identified in Figure 2).

Known modifications include, but are not limited to, acetylation, acylation, ADP-ribosylation, amidation, covalent attachment of flavin, covalent attachment of a heme moiety, covalent attachment of a nucleotide or nucleotide derivative, covalent attachment of a lipid or lipid derivative, covalent attachment of phosphotidylinositol, cross-linking, cyclization, disulfide bond formation, demethylation, formation of covalent crosslinks, formation of cystine, formation of pyroglutamate, formylation, gamma carboxylation, glycosylation, GPI anchor formation, hydroxylation, iodination, methylation, myristoylation, oxidation, proteolytic processing, phosphorylation, prenylation, racemization, selenoylation, sulfation, transfer-RNA mediated addition of amino acids to proteins such as arginylation, and ubiquitination.

Such modifications are well known to those of skill in the art and have been described in great detail in the scientific literature. Several particularly common modifications, glycosylation, lipid attachment, sulfation, gamma-carboxylation of glutamic acid residues, hydroxylation and ADP-ribosylation, for instance, are described in most basic texts, such as *Proteins - Structure and Molecular Properties*, 2nd Ed., T.E. Creighton, W. H. Freeman and Company, New York (1993). Many detailed reviews are available on this subject, such as by Wold, F., *Posttranslational Covalent Modification of Proteins*, B.C. Johnson, Ed., Academic Press, New York 1-12 (1983); Seifter *et al.* (*Meth. Enzymol. 182*: 626-646 (1990)) and Rattan *et al.* (*Ann. N.Y. Acad. Sci. 663*:48-62 (1992)).

Accordingly, the transporter peptides of the present invention also encompass derivatives or analogs in which a substituted amino acid residue is not one encoded by the genetic code, in which a substituent group is included, in which the mature transporter peptide is fused with another compound, such as a compound to increase the half-life of the transporter peptide (for example, polyethylene glycol), or in which the additional amino acids are fused to the mature transporter peptide, such as a leader or secretory sequence or a sequence for purification of the mature transporter peptide or a pro-protein sequence.

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#### Protein/Peptide Uses

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The proteins of the present invention can be used in substantial and specific assays related to the functional information provided in the Figures; to raise antibodies or to elicit another immune response; as a reagent (including the labeled reagent) in assays designed to quantitatively determine levels of the protein (or its binding partner or ligand) in biological fluids; and as markers for tissues in which the corresponding protein is preferentially expressed (either constitutively or at a particular stage of tissue differentiation or development or in a disease state). Where the protein binds or potentially binds to another protein or ligand (such as, for example, in a transporter-effector protein interaction or transporter-ligand interaction), the protein can be used to identify the binding partner/ligand so as to develop a system to identify inhibitors of the binding interaction. Any or all of these uses are capable of being developed into reagent grade or kit format for commercialization as commercial products.

Methods for performing the uses listed above are well known to those skilled in the art. References disclosing such methods include "Molecular Cloning: A Laboratory Manual", 2d ed., Cold Spring Harbor Laboratory Press, Sambrook, J., E. F. Fritsch and T. Maniatis eds., 1989, and "Methods in Enzymology: Guide to Molecular Cloning Techniques", Academic Press, Berger, S. L. and A. R. Kimmel eds., 1987.

The potential uses of the peptides of the present invention are based primarily on the source of the protein as well as the class/action of the protein. For example, transporters isolated from humans and their human/mammalian orthologs serve as targets for identifying agents for use in mammalian therapeutic applications, e.g. a human drug, particularly in modulating a biological or pathological response in a cell or tissue that expresses the transporter.

Experimental data as provided in Figure 1 indicates that sodium/calcium exchanger proteins of the present invention are expressed in humans in the heart, retina, kidney, fetal brain, and fetal heart. Specifically, a virtual northern blot shows expression in the fetal brain. In addition, PCR-based tissue screening panel indicates expression in brain, heart, kidney, lung, spleen, testis, leukocyte and fetal brain. A large percentage of pharmaceutical agents are being developed that modulate the activity of transporter proteins, particularly members of the sodium/calcium exchanger subfamily (see Background of the Invention). The structural and functional information provided in the Background and Figures provide specific and substantial uses for the molecules of the present invention, particularly in combination with the expression information provided in Figure 1. Experimental data as provided in Figure 1 indicates expression in humans in

brain, heart, kidney, lung, spleen, testis, leukocyte and fetal brain. Such uses can readily be determined using the information provided herein, that known in the art and routine experimentation.

The proteins of the present invention (including variants and fragments that may have been disclosed prior to the present invention) are useful for biological assays related to transporters that are related to members of the sodium/calcium exchanger subfamily. Such assays involve any of the known transporter functions or activities or properties useful for diagnosis and treatment of transporter-related conditions that are specific for the subfamily of transporters that the one of the present invention belongs to, particularly in cells and tissues that express the transporter.

Experimental data as provided in Figure 1 indicates that sodium/calcium exchanger proteins of the present invention are expressed in humans in the heart, retina, kidney, fetal brain, and fetal heart. Specifically, a virtual northern blot shows expression in the fetal brain. In addition, PCR-based tissue screening panel indicates expression in brain, heart, kidney, lung, spleen, testis, leukocyte and fetal brain.

The proteins of the present invention are also useful in drug screening assays, in cell-based or cell-free systems ((Hodgson, Bio/technology, 1992, Sept 10(9);973-80). Cell-based systems can be native, i.e., cells that normally express the transporter, as a biopsy or expanded in cell culture. Experimental data as provided in Figure 1 indicates expression in humans in brain, heart, kidney, lung, spleen, testis, leukocyte and fetal brain. In an alternate embodiment, cell-based assays involve recombinant host cells expressing the transporter protein.

The polypeptides can be used to identify compounds that modulate transporter activity of the protein in its natural state or an altered form that causes a specific disease or pathology associated with the transporter. Both the transporters of the present invention and appropriate variants and fragments can be used in high-throughput screens to assay candidate compounds for the ability to bind to the transporter. These compounds can be further screened against a functional transporter to determine the effect of the compound on the transporter activity. Further, these compounds can be tested in animal or invertebrate systems to determine activity/effectiveness. Compounds can be identified that activate (agonist) or inactivate (antagonist) the transporter to a desired degree.

Further, the proteins of the present invention can be used to screen a compound for the ability to stimulate or inhibit interaction between the transporter protein and a molecule that normally interacts with the transporter protein, e.g. a substrate or a component of the signal pathway that the transporter protein normally interacts (for example, another transporter). Such assays

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typically include the steps of combining the transporter protein with a candidate compound under conditions that allow the transporter protein, or fragment, to interact with the target molecule, and to detect the formation of a complex between the protein and the target or to detect the biochemical consequence of the interaction with the transporter protein and the target, such as any of the associated effects of signal transduction such as changes in membrane potential, protein phosphorylation, cAMP turnover, and adenylate cyclase activation, etc.

Candidate compounds include, for example, 1) peptides such as soluble peptides, including Ig-tailed fusion peptides and members of random peptide libraries (see, e.g., Lam et al., Nature 354:82-84 (1991); Houghten et al., Nature 354:84-86 (1991)) and combinatorial chemistry-derived molecular libraries made of D- and/or L- configuration amino acids; 2) phosphopeptides (e.g., members of random and partially degenerate, directed phosphopeptide libraries, see, e.g., Songyang et al., Cell 72:767-778 (1993)); 3) antibodies (e.g., polyclonal, monoclonal, humanized, anti-idiotypic, chimeric, and single chain antibodies as well as Fab, F(ab')<sub>2</sub>, Fab expression library fragments, and epitope-binding fragments of antibodies); and 4) small organic and inorganic molecules (e.g., molecules obtained from combinatorial and natural product libraries).

One candidate compound is a soluble fragment of the receptor that competes for ligand binding. Other candidate compounds include mutant transporters or appropriate fragments containing mutations that affect transporter function and thus compete for ligand. Accordingly, a fragment that competes for ligand, for example with a higher affinity, or a fragment that binds ligand but does not allow release, is encompassed by the invention.

The invention further includes other end point assays to identify compounds that modulate (stimulate or inhibit) transporter activity. The assays typically involve an assay of events in the signal transduction pathway that indicate transporter activity. Thus, the transport of a ligand, change in cell membrane potential, activation of a protein, a change in the expression of genes that are up- or down-regulated in response to the transporter protein dependent signal cascade can be assayed.

Any of the biological or biochemical functions mediated by the transporter can be used as an endpoint assay. These include all of the biochemical or biochemical/biological events described herein, in the references cited herein, incorporated by reference for these endpoint assay targets, and other functions known to those of ordinary skill in the art or that can be readily identified using the information provided in the Figures, particularly Figure 2. Specifically, a biological function of a cell or tissues that expresses the transporter can be assayed. Experimental data as provided in Figure 1 indicates that sodium/calcium exchanger proteins of the present invention are expressed in

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humans in the heart, retina, kidney, fetal brain, and fetal heart. Specifically, a virtual northern blot shows expression in the fetal brain. In addition, PCR-based tissue screening panel indicates expression in brain, heart, kidney, lung, spleen, testis, leukocyte and fetal brain.

Binding and/or activating compounds can also be screened by using chimeric transporter proteins in which the amino terminal extracellular domain, or parts thereof, the entire transmembrane domain or subregions, such as any of the seven transmembrane segments or any of the intracellular or extracellular loops and the carboxy terminal intracellular domain, or parts thereof, can be replaced by heterologous domains or subregions. For example, a ligand-binding region can be used that interacts with a different ligand then that which is recognized by the native transporter. Accordingly, a different set of signal transduction components is available as an endpoint assay for activation. This allows for assays to be performed in other than the specific host cell from which the transporter is derived.

The proteins of the present invention are also useful in competition binding assays in methods designed to discover compounds that interact with the transporter (e.g. binding partners and/or ligands). Thus, a compound is exposed to a transporter polypeptide under conditions that allow the compound to bind or to otherwise interact with the polypeptide. Soluble transporter polypeptide is also added to the mixture. If the test compound interacts with the soluble transporter polypeptide, it decreases the amount of complex formed or activity from the transporter target. This type of assay is particularly useful in cases in which compounds are sought that interact with specific regions of the transporter. Thus, the soluble polypeptide that competes with the target transporter region is designed to contain peptide sequences corresponding to the region of interest.

To perform cell free drug screening assays, it is sometimes desirable to immobilize either the transporter protein, or fragment, or its target molecule to facilitate separation of complexes from uncomplexed forms of one or both of the proteins, as well as to accommodate automation of the assay.

Techniques for immobilizing proteins on matrices can be used in the drug screening assays. In one embodiment, a fusion protein can be provided which adds a domain that allows the protein to be bound to a matrix. For example, glutathione-S-transferase fusion proteins can be adsorbed onto glutathione sepharose beads (Sigma Chemical, St. Louis, MO) or glutathione derivatized microtitre plates, which are then combined with the cell lysates (e.g., <sup>35</sup>S-labeled) and the candidate compound, and the mixture incubated under conditions conducive to complex formation (e.g., at physiological conditions for salt and pH). Following incubation, the beads are washed to remove any unbound label, and the matrix immobilized and radiolabel determined directly, or in the

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supernatant after the complexes are dissociated. Alternatively, the complexes can be dissociated from the matrix, separated by SDS-PAGE, and the level of transporter-binding protein found in the bead fraction quantitated from the gel using standard electrophoretic techniques. For example, either the polypeptide or its target molecule can be immobilized utilizing conjugation of biotin and streptavidin using techniques well known in the art. Alternatively, antibodies reactive with the protein but which do not interfere with binding of the protein to its target molecule can be derivatized to the wells of the plate, and the protein trapped in the wells by antibody conjugation. Preparations of a transporter-binding protein and a candidate compound are incubated in the transporter protein-presenting wells and the amount of complex trapped in the well can be quantitated. Methods for detecting such complexes, in addition to those described above for the GST-immobilized complexes, include immunodetection of complexes using antibodies reactive with the transporter protein target molecule, or which are reactive with transporter protein and compete with the target molecule, as well as enzyme-linked assays which rely on detecting an enzymatic activity associated with the target molecule.

Agents that modulate one of the transporters of the present invention can be identified using one or more of the above assays, alone or in combination. It is generally preferable to use a cell-based or cell free system first and then confirm activity in an animal or other model system. Such model systems are well known in the art and can readily be employed in this context.

Modulators of transporter protein activity identified according to these drug screening assays can be used to treat a subject with a disorder mediated by the transporter pathway, by treating cells or tissues that express the transporter. Experimental data as provided in Figure 1 indicates expression in humans in brain, heart, kidney, lung, spleen, testis, leukocyte and fetal brain. These methods of treatment include the steps of administering a modulator of transporter activity in a pharmaceutical composition to a subject in need of such treatment, the modulator being identified as described herein.

In yet another aspect of the invention, the transporter proteins can be used as "bait proteins" in a two-hybrid assay or three-hybrid assay (see, e.g., U.S. Patent No. 5,283,317; Zervos et al. (1993) Cell 72:223-232; Madura et al. (1993) J. Biol. Chem. 268:12046-12054; Bartel et al. (1993) Biotechniques 14:920-924; Iwabuchi et al. (1993) Oncogene 8:1693-1696; and Brent WO94/10300), to identify other proteins, which bind to or interact with the transporter and are involved in transporter activity. Such transporter-binding proteins are also likely to be involved in the propagation of signals by the transporter proteins or transporter targets as, for

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example, downstream elements of a transporter-mediated signaling pathway. Alternatively, such transporter-binding proteins are likely to be transporter inhibitors.

The two-hybrid system is based on the modular nature of most transcription factors, which consist of separable DNA-binding and activation domains. Briefly, the assay utilizes two different DNA constructs. In one construct, the gene that codes for a transporter protein is fused to a gene encoding the DNA binding domain of a known transcription factor (e.g., GAL-4). In the other construct, a DNA sequence, from a library of DNA sequences, that encodes an unidentified protein ("prey" or "sample") is fused to a gene that codes for the activation domain of the known transcription factor. If the "bait" and the "prey" proteins are able to interact, *in vivo*, forming a transporter-dependent complex, the DNA-binding and activation domains of the transcription factor are brought into close proximity. This proximity allows transcription of a reporter gene (e.g., LacZ) which is operably linked to a transcriptional regulatory site responsive to the transcription factor. Expression of the reporter gene can be detected and cell colonies containing the functional transcription factor can be isolated and used to obtain the cloned gene which encodes the protein which interacts with the transporter protein.

This invention further pertains to novel agents identified by the above-described screening assays. Accordingly, it is within the scope of this invention to further use an agent identified as described herein in an appropriate animal model. For example, an agent identified as described herein (e.g., a transporter-modulating agent, an antisense transporter nucleic acid molecule, a transporter-specific antibody, or a transporter-binding partner) can be used in an animal or other model to determine the efficacy, toxicity, or side effects of treatment with such an agent. Alternatively, an agent identified as described herein can be used in an animal or other model to determine the mechanism of action of such an agent. Furthermore, this invention pertains to uses of novel agents identified by the above-described screening assays for treatments as described herein.

The transporter proteins of the present invention are also useful to provide a target for diagnosing a disease or predisposition to disease mediated by the peptide. Accordingly, the invention provides methods for detecting the presence, or levels of, the protein (or encoding mRNA) in a cell, tissue, or organism. Experimental data as provided in Figure 1 indicates expression in humans in brain, heart, kidney, lung, spleen, testis, leukocyte and fetal brain. The method involves contacting a biological sample with a compound capable of interacting with the transporter protein such that the interaction can be detected. Such an assay can be provided in a single detection format or a multi-detection format such as an antibody chip array.

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One agent for detecting a protein in a sample is an antibody capable of selectively binding to protein. A biological sample includes tissues, cells and biological fluids isolated from a subject, as well as tissues, cells and fluids present within a subject.

The peptides of the present invention also provide targets for diagnosing active protein activity, disease, or predisposition to disease, in a patient having a variant peptide, particularly activities and conditions that are known for other members of the family of proteins to which the present one belongs. Thus, the peptide can be isolated from a biological sample and assayed for the presence of a genetic mutation that results in aberrant peptide. This includes amino acid substitution, deletion, insertion, rearrangement, (as the result of aberrant splicing events), and inappropriate post-translational modification. Analytic methods include altered electrophoretic mobility, altered tryptic peptide digest, altered transporter activity in cell-based or cell-free assay, alteration in ligand or antibody-binding pattern, altered isoelectric point, direct amino acid sequencing, and any other of the known assay techniques useful for detecting mutations in a protein. Such an assay can be provided in a single detection format or a multi-detection format such as an antibody chip array.

In vitro techniques for detection of peptide include enzyme linked immunosorbent assays (ELISAs), Western blots, immunoprecipitations and immunofluorescence using a detection reagent, such as an antibody or protein binding agent. Alternatively, the peptide can be detected in vivo in a subject by introducing into the subject a labeled anti-peptide antibody or other types of detection agent. For example, the antibody can be labeled with a radioactive marker whose presence and location in a subject can be detected by standard imaging techniques. Particularly useful are methods that detect the allelic variant of a peptide expressed in a subject and methods which detect fragments of a peptide in a sample.

The peptides are also useful in pharmacogenomic analysis. Pharmacogenomics deal with clinically significant hereditary variations in the response to drugs due to altered drug disposition and abnormal action in affected persons. See, e.g., Eichelbaum, M. (Clin. Exp. Pharmacol. Physiol. 23(10-11):983-985 (1996)), and Linder, M.W. (Clin. Chem. 43(2):254-266 (1997)). The clinical outcomes of these variations result in severe toxicity of therapeutic drugs in certain individuals or therapeutic failure of drugs in certain individuals as a result of individual variation in metabolism. Thus, the genotype of the individual can determine the way a therapeutic compound acts on the body or the way the body metabolizes the compound. Further, the activity of drug metabolizing enzymes effects both the intensity and duration of drug action. Thus, the pharmacogenomics of the individual permit the selection of effective compounds and effective dosages of such compounds for

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prophylactic or therapeutic treatment based on the individual's genotype. The discovery of genetic polymorphisms in some drug metabolizing enzymes has explained why some patients do not obtain the expected drug effects, show an exaggerated drug effect, or experience serious toxicity from standard drug dosages. Polymorphisms can be expressed in the phenotype of the extensive metabolizer and the phenotype of the poor metabolizer. Accordingly, genetic polymorphism may lead to allelic protein variants of the transporter protein in which one or more of the transporter functions in one population is different from those in another population. The peptides thus allow a target to ascertain a genetic predisposition that can affect treatment modality. Thus, in a ligand-based treatment, polymorphism may give rise to amino terminal extracellular domains and/or other ligand-binding regions that are more or less active in ligand binding, and transporter activation. Accordingly, ligand dosage would necessarily be modified to maximize the therapeutic effect within a given population containing a polymorphism. As an alternative to genotyping, specific polymorphic peptides could be identified.

The peptides are also useful for treating a disorder characterized by an absence of, inappropriate, or unwanted expression of the protein. Experimental data as provided in Figure 1 indicates expression in humans in brain, heart, kidney, lung, spleen, testis, leukocyte and fetal brain. Accordingly, methods for treatment include the use of the transporter protein or fragments.

#### **Antibodies**

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The invention also provides antibodies that selectively bind to one of the peptides of the present invention, a protein comprising such a peptide, as well as variants and fragments thereof. As used herein, an antibody selectively binds a target peptide when it binds the target peptide and does not significantly bind to unrelated proteins. An antibody is still considered to selectively bind a peptide even if it also binds to other proteins that are not substantially homologous with the target peptide so long as such proteins share homology with a fragment or domain of the peptide target of the antibody. In this case, it would be understood that antibody binding to the peptide is still selective despite some degree of cross-reactivity.

As used herein, an antibody is defined in terms consistent with that recognized within the art: they are multi-subunit proteins produced by a mammalian organism in response to an antigen challenge. The antibodies of the present invention include polyclonal antibodies and monoclonal antibodies, as well as fragments of such antibodies, including, but not limited to, Fab or F(ab')<sub>2</sub>, and Fv fragments.

Many methods are known for generating and/or identifying antibodies to a given target peptide. Several such methods are described by Harlow, Antibodies, Cold Spring Harbor Press, (1989).

In general, to generate antibodies, an isolated peptide is used as an immunogen and is administered to a mammalian organism, such as a rat, rabbit or mouse. The full-length protein, an antigenic peptide fragment or a fusion protein can be used. Particularly important fragments are those covering functional domains, such as the domains identified in Figure 2, and domain of sequence homology or divergence amongst the family, such as those that can readily be identified using protein alignment methods and as presented in the Figures.

Antibodies are preferably prepared from regions or discrete fragments of the transporter proteins. Antibodies can be prepared from any region of the peptide as described herein. However, preferred regions will include those involved in function/activity and/or transporter/binding partner interaction. Figure 2 can be used to identify particularly important regions while sequence alignment can be used to identify conserved and unique sequence fragments.

An antigenic fragment will typically comprise at least 8 contiguous amino acid residues. The antigenic peptide can comprise, however, at least 10, 12, 14, 16 or more amino acid residues. Such fragments can be selected on a physical property, such as fragments correspond to regions that are located on the surface of the protein, e.g., hydrophilic regions or can be selected based on sequence uniqueness (see Figure 2).

Detection on an antibody of the present invention can be facilitated by coupling (i.e., physically linking) the antibody to a detectable substance. Examples of detectable substances include various enzymes, prosthetic groups, fluorescent materials, luminescent materials, bioluminescent materials, and radioactive materials. Examples of suitable enzymes include horseradish peroxidase, alkaline phosphatase, β-galactosidase, or acetylcholinesterase; examples of suitable prosthetic group complexes include streptavidin/biotin and avidin/biotin; examples of suitable fluorescent materials include umbelliferone, fluorescein, fluorescein isothiocyanate, rhodamine, dichlorotriazinylamine fluorescein, dansyl chloride or phycoerythrin; an example of a luminescent material includes luminol; examples of bioluminescent materials include luciferase, luciferin, and aequorin, and examples of suitable radioactive material include <sup>125</sup>I, <sup>131</sup>I, <sup>35</sup>S or <sup>3</sup>H.

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#### Antibody Uses

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The antibodies can be used to isolate one of the proteins of the present invention by standard techniques, such as affinity chromatography or immunoprecipitation. The antibodies can facilitate the purification of the natural protein from cells and recombinantly produced protein expressed in host cells. In addition, such antibodies are useful to detect the presence of one of the proteins of the present invention in cells or tissues to determine the pattern of expression of the protein among various tissues in an organism and over the course of normal development. Experimental data as provided in Figure 1 indicates that sodium/calcium exchanger proteins of the present invention are expressed in humans in the heart, retina, kidney, fetal brain, and fetal heart. Specifically, a virtual northern blot shows expression in the fetal brain. In addition, PCR-based tissue screening panel indicates expression in brain, heart, kidney, lung, spleen, testis, leukocyte and fetal brain. Further, such antibodies can be used to detect protein *in situ*, *in vitro*, or in a cell lysate or supernatant in order to evaluate the abundance and pattern of expression. Also, such antibodies can be used to assess abnormal tissue distribution or abnormal expression during development or progression of a biological condition. Antibody detection of circulating fragments of the full length protein can be used to identify turnover.

Further, the antibodies can be used to assess expression in disease states such as in active stages of the disease or in an individual with a predisposition toward disease related to the protein's function. When a disorder is caused by an inappropriate tissue distribution, developmental expression, level of expression of the protein, or expressed/processed form, the antibody can be prepared against the normal protein. Experimental data as provided in Figure 1 indicates expression in humans in brain, heart, kidney, lung, spleen, testis, leukocyte and fetal brain. If a disorder is characterized by a specific mutation in the protein, antibodies specific for this mutant protein can be used to assay for the presence of the specific mutant protein.

The antibodies can also be used to assess normal and aberrant subcellular localization of cells in the various tissues in an organism. Experimental data as provided in Figure 1 indicates expression in humans in brain, heart, kidney, lung, spleen, testis, leukocyte and fetal brain. The diagnostic uses can be applied, not only in genetic testing, but also in monitoring a treatment modality. Accordingly, where treatment is ultimately aimed at correcting expression level or the presence of aberrant sequence and aberrant tissue distribution or developmental expression, antibodies directed against the protein or relevant fragments can be used to monitor therapeutic efficacy.

Additionally, antibodies are useful in pharmacogenomic analysis. Thus, antibodies prepared against polymorphic proteins can be used to identify individuals that require modified treatment modalities. The antibodies are also useful as diagnostic tools as an immunological marker for aberrant protein analyzed by electrophoretic mobility, isoelectric point, tryptic peptide digest, and other physical assays known to those in the art.

The antibodies are also useful for tissue typing. Experimental data as provided in Figure 1 indicates expression in humans in brain, heart, kidney, lung, spleen, testis, leukocyte and fetal brain. Thus, where a specific protein has been correlated with expression in a specific tissue, antibodies that are specific for this protein can be used to identify a tissue type.

The antibodies are also useful for inhibiting protein function, for example, blocking the binding of the transporter peptide to a binding partner such as a ligand or protein binding partner. These uses can also be applied in a therapeutic context in which treatment involves inhibiting the protein's function. An antibody can be used, for example, to block binding, thus modulating (agonizing or antagonizing) the peptides activity. Antibodies can be prepared against specific fragments containing sites required for function or against intact protein that is associated with a cell or cell membrane. See Figure 2 for structural information relating to the proteins of the present invention.

The invention also encompasses kits for using antibodies to detect the presence of a protein in a biological sample. The kit can comprise antibodies such as a labeled or labelable antibody and a compound or agent for detecting protein in a biological sample; means for determining the amount of protein in the sample; means for comparing the amount of protein in the sample with a standard; and instructions for use. Such a kit can be supplied to detect a single protein or epitope or can be configured to detect one of a multitude of epitopes, such as in an antibody detection array. Arrays are described in detail below for nucleic acid arrays and similar methods have been developed for antibody arrays.

#### Nucleic Acid Molecules

The present invention further provides isolated nucleic acid molecules that encode a transporter peptide or protein of the present invention (cDNA, transcript and genomic sequence). Such nucleic acid molecules will consist of, consist essentially of, or comprise a nucleotide sequence that encodes one of the transporter peptides of the present invention, an allelic variant thereof, or an ortholog or paralog thereof.

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As used herein, an "isolated" nucleic acid molecule is one that is separated from other nucleic acid present in the natural source of the nucleic acid. Preferably, an "isolated" nucleic acid is free of sequences that naturally flank the nucleic acid (i.e., sequences located at the 5' and 3' ends of the nucleic acid) in the genomic DNA of the organism from which the nucleic acid is derived. However, there can be some flanking nucleotide sequences, for example up to about 5KB, 4KB, 3KB, 2KB, or 1KB or less, particularly contiguous peptide encoding sequences and peptide encoding sequences within the same gene but separated by introns in the genomic sequence. The important point is that the nucleic acid is isolated from remote and unimportant flanking sequences such that it can be subjected to the specific manipulations described herein such as recombinant expression, preparation of probes and primers, and other uses specific to the nucleic acid sequences.

Moreover, an "isolated" nucleic acid molecule, such as a transcript/cDNA molecule, can be substantially free of other cellular material, or culture medium when produced by recombinant techniques, or chemical precursors or other chemicals when chemically synthesized. However, the nucleic acid molecule can be fused to other coding or regulatory sequences and still be considered isolated.

For example, recombinant DNA molecules contained in a vector are considered isolated. Further examples of isolated DNA molecules include recombinant DNA molecules maintained in heterologous host cells or purified (partially or substantially) DNA molecules in solution. Isolated RNA molecules include *in vivo* or *in vitro* RNA transcripts of the isolated DNA molecules of the present invention. Isolated nucleic acid molecules according to the present invention further include such molecules produced synthetically.

Accordingly, the present invention provides nucleic acid molecules that consist of the nucleotide sequence shown in Figure 1 or 3 (SEQ ID NO:1, transcript sequence and SEQ ID NO:3, genomic sequence), or any nucleic acid molecule that encodes the protein provided in Figure 2, SEQ ID NO:2. A nucleic acid molecule consists of a nucleotide sequence when the nucleotide sequence is the complete nucleotide sequence of the nucleic acid molecule.

The present invention further provides nucleic acid molecules that consist essentially of the nucleotide sequence shown in Figure 1 or 3 (SEQ ID NO:1, transcript sequence and SEQ ID NO:3, genomic sequence), or any nucleic acid molecule that encodes the protein provided in Figure 2, SEQ ID NO:2. A nucleic acid molecule consists essentially of a nucleotide sequence when such a nucleotide sequence is present with only a few additional nucleic acid residues in the final nucleic acid molecule.

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The present invention further provides nucleic acid molecules that comprise the nucleotide sequences shown in Figure 1 or 3 (SEQ ID NO:1, transcript sequence and SEQ ID NO:3, genomic sequence), or any nucleic acid molecule that encodes the protein provided in Figure 2, SEQ ID NO:2. A nucleic acid molecule comprises a nucleotide sequence when the nucleotide sequence is at least part of the final nucleotide sequence of the nucleic acid molecule. In such a fashion, the nucleic acid molecule can be only the nucleotide sequence or have additional nucleic acid residues, such as nucleic acid residues that are naturally associated with it or heterologous nucleotide sequences. Such a nucleic acid molecule can have a few additional nucleotides or can comprise several hundred or more additional nucleotides. A brief description of how various types of these nucleic acid molecules can be readily made/isolated is provided below.

In Figures 1 and 3, both coding and non-coding sequences are provided. Because of the source of the present invention, humans genomic sequence (Figure 3) and cDNA/transcript sequences (Figure 1), the nucleic acid molecules in the Figures will contain genomic intronic sequences, 5' and 3' non-coding sequences, gene regulatory regions and non-coding intergenic sequences. In general such sequence features are either noted in Figures 1 and 3 or can readily be identified using computational tools known in the art. As discussed below, some of the non-coding regions, particularly gene regulatory elements such as promoters, are useful for a variety of purposes, e.g. control of heterologous gene expression, target for identifying gene activity modulating compounds, and are particularly claimed as fragments of the genomic sequence provided herein.

The isolated nucleic acid molecules can encode the mature protein plus additional amino or carboxyl-terminal amino acids, or amino acids interior to the mature peptide (when the mature form has more than one peptide chain, for instance). Such sequences may play a role in processing of a protein from precursor to a mature form, facilitate protein trafficking, prolong or shorten protein half-life or facilitate manipulation of a protein for assay or production, among other things. As generally is the case *in situ*, the additional amino acids may be processed away from the mature protein by cellular enzymes.

As mentioned above, the isolated nucleic acid molecules include, but are not limited to, the sequence encoding the transporter peptide alone, the sequence encoding the mature peptide and additional coding sequences, such as a leader or secretory sequence (e.g., a pre-pro or pro-protein sequence), the sequence encoding the mature peptide, with or without the additional coding sequences, plus additional non-coding sequences, for example introns and non-coding 5' and 3' sequences such as transcribed but non-translated sequences that play a role in transcription, mRNA

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processing (including splicing and polyadenylation signals), ribosome binding and stability of mRNA. In addition, the nucleic acid molecule may be fused to a marker sequence encoding, for example, a peptide that facilitates purification.

Isolated nucleic acid molecules can be in the form of RNA, such as mRNA, or in the form DNA, including cDNA and genomic DNA obtained by cloning or produced by chemical synthetic techniques or by a combination thereof. The nucleic acid, especially DNA, can be double-stranded or single-stranded. Single-stranded nucleic acid can be the coding strand (sense strand) or the non-coding strand (anti-sense strand).

The invention further provides nucleic acid molecules that encode fragments of the peptides of the present invention as well as nucleic acid molecules that encode obvious variants of the transporter proteins of the present invention that are described above. Such nucleic acid molecules may be naturally occurring, such as allelic variants (same locus), paralogs (different locus), and orthologs (different organism), or may be constructed by recombinant DNA methods or by chemical synthesis. Such non-naturally occurring variants may be made by mutagenesis techniques, including those applied to nucleic acid molecules, cells, or organisms. Accordingly, as discussed above, the variants can contain nucleotide substitutions, deletions, inversions and insertions. Variation can occur in either or both the coding and non-coding regions. The variations can produce both conservative and non-conservative amino acid substitutions.

The present invention further provides non-coding fragments of the nucleic acid molecules provided in Figures 1 and 3. Preferred non-coding fragments include, but are not limited to, promoter sequences, enhancer sequences, gene modulating sequences and gene termination sequences. Such fragments are useful in controlling heterologous gene expression and in developing screens to identify gene-modulating agents. A promoter can readily be identified as being 5' to the ATG start site in the genomic sequence provided in Figure 3.

A fragment comprises a contiguous nucleotide sequence greater than 12 or more nucleotides. Further, a fragment could at least 30, 40, 50, 100, 250 or 500 nucleotides in length. The length of the fragment will be based on its intended use. For example, the fragment can encode epitope bearing regions of the peptide, or can be useful as DNA probes and primers. Such fragments can be isolated using the known nucleotide sequence to synthesize an oligonucleotide probe. A labeled probe can then be used to screen a cDNA library, genomic DNA library, or mRNA to isolate nucleic acid corresponding to the coding region. Further, primers can be used in PCR reactions to clone specific regions of gene.

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A probe/primer typically comprises substantially a purified oligonucleotide or oligonucleotide pair. The oligonucleotide typically comprises a region of nucleotide sequence that hybridizes under stringent conditions to at least about 12, 20, 25, 40, 50 or more consecutive nucleotides.

Orthologs, homologs, and allelic variants can be identified using methods well known in the art. As described in the Peptide Section, these variants comprise a nucleotide sequence encoding a peptide that is typically 60-70%, 70-80%, 80-90%, and more typically at least about 90-95% or more homologous to the nucleotide sequence shown in the Figure sheets or a fragment of this sequence. Such nucleic acid molecules can readily be identified as being able to hybridize under moderate to stringent conditions, to the nucleotide sequence shown in the Figure sheets or a fragment of the sequence. Allelic variants can readily be determined by genetic locus of the encoding gene. As indicated by the data presented in Figure 3, the map position was determined to be on chromosome 14 by ePCR.

Figure 3 provides information on SNPs that have been identified in a gene encoding the transporter protein of the present invention. 140 SNP variants were found, including 6 indels (indicated by a "-") and 1 SNPs in exons. The others were found in in introns and regions 5' and 3' of the ORF. Such SNPs in introns and outside the ORF may affect control/regulatory elements.

As used herein, the term "hybridizes under stringent conditions" is intended to describe conditions for hybridization and washing under which nucleotide sequences encoding a peptide at least 60-70% homologous to each other typically remain hybridized to each other. The conditions can be such that sequences at least about 60%, at least about 70%, or at least about 80% or more homologous to each other typically remain hybridized to each other. Such stringent conditions are known to those skilled in the art and can be found in *Current Protocols in Molecular Biology*, John Wiley & Sons, N.Y. (1989), 6.3.1-6.3.6. One example of stringent hybridization conditions are hybridization in 6X sodium chloride/sodium citrate (SSC) at about 45C, followed by one or more washes in 0.2 X SSC, 0.1% SDS at 50-65C. Examples of moderate to low stringency hybridization conditions are well known in the art.

### Nucleic Acid Molecule Uses

The nucleic acid molecules of the present invention are useful for probes, primers, chemical intermediates, and in biological assays. The nucleic acid molecules are useful as a hybridization probe for messenger RNA, transcript/cDNA and genomic DNA to isolate full-length cDNA and

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genomic clones encoding the peptide described in Figure 2 and to isolate cDNA and genomic clones that correspond to variants (alleles, orthologs, etc.) producing the same or related peptides shown in Figure 2. 140 SNPs, including 6 indels, have been identified in the gene encoding the transporter protein provided by the present invention and are given in Figure 3.

The probe can correspond to any sequence along the entire length of the nucleic acid molecules provided in the Figures. Accordingly, it could be derived from 5' noncoding regions, the coding region, and 3' noncoding regions. However, as discussed, fragments are not to be construed as encompassing fragments disclosed prior to the present invention.

The nucleic acid molecules are also useful as primers for PCR to amplify any given region of a nucleic acid molecule and are useful to synthesize antisense molecules of desired length and sequence.

The nucleic acid molecules are also useful for constructing recombinant vectors. Such vectors include expression vectors that express a portion of, or all of, the peptide sequences. Vectors also include insertion vectors, used to integrate into another nucleic acid molecule sequence, such as into the cellular genome, to alter *in situ* expression of a gene and/or gene product. For example, an endogenous coding sequence can be replaced via homologous recombination with all or part of the coding region containing one or more specifically introduced mutations.

The nucleic acid molecules are also useful for expressing antigenic portions of the proteins.

The nucleic acid molecules are also useful as probes for determining the chromosomal positions of the nucleic acid molecules by means of *in situ* hybridization methods. As indicated by the data presented in Figure 3, the map position was determined to be on chromosome 14 by ePCR.

The nucleic acid molecules are also useful in making vectors containing the gene regulatory regions of the nucleic acid molecules of the present invention.

The nucleic acid molecules are also useful for designing ribozymes corresponding to all, or a part, of the mRNA produced from the nucleic acid molecules described herein.

The nucleic acid molecules are also useful for making vectors that express part, or all, of the peptides.

The nucleic acid molecules are also useful for constructing host cells expressing a part, or all, of the nucleic acid molecules and peptides.

The nucleic acid molecules are also useful for constructing transgenic animals expressing all, or a part, of the nucleic acid molecules and peptides.

The nucleic acid molecules are also useful as hybridization probes for determining the presence, level, form and distribution of nucleic acid expression. Experimental data as provided in

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Figure 1 indicates that sodium/calcium exchanger proteins of the present invention are expressed in humans in the heart, retina, kidney, fetal brain, and fetal heart. Specifically, a virtual northern blot shows expression in the fetal brain. In addition, PCR-based tissue screening panel indicates expression in brain, heart, kidney, lung, spleen, testis, leukocyte and fetal brain.

Accordingly, the probes can be used to detect the presence of, or to determine levels of, a specific nucleic acid molecule in cells, tissues, and in organisms. The nucleic acid whose level is determined can be DNA or RNA. Accordingly, probes corresponding to the peptides described herein can be used to assess expression and/or gene copy number in a given cell, tissue, or organism. These uses are relevant for diagnosis of disorders involving an increase or decrease in transporter protein expression relative to normal results.

In vitro techniques for detection of mRNA include Northern hybridizations and in situ hybridizations. In vitro techniques for detecting DNA include Southern hybridizations and in situ hybridization.

Probes can be used as a part of a diagnostic test kit for identifying cells or tissues that express a transporter protein, such as by measuring a level of a transporter-encoding nucleic acid in a sample of cells from a subject e.g., mRNA or genomic DNA, or determining if a transporter gene has been mutated. Experimental data as provided in Figure 1 indicates that sodium/calcium exchanger proteins of the present invention are expressed in humans in the heart, retina, kidney, fetal brain, and fetal heart. Specifically, a virtual northern blot shows expression in the fetal brain. In addition, PCR-based tissue screening panel indicates expression in brain, heart, kidney, lung, spleen, testis, leukocyte and fetal brain.

Nucleic acid expression assays are useful for drug screening to identify compounds that modulate transporter nucleic acid expression.

The invention thus provides a method for identifying a compound that can be used to treat a disorder associated with nucleic acid expression of the transporter gene, particularly biological and pathological processes that are mediated by the transporter in cells and tissues that express it.

Experimental data as provided in Figure 1 indicates expression in humans in brain, heart, kidney, lung, spleen, testis, leukocyte and fetal brain. The method typically includes assaying the ability of the compound to modulate the expression of the transporter nucleic acid and thus identifying a compound that can be used to treat a disorder characterized by undesired transporter nucleic acid expression. The assays can be performed in cell-based and cell-free systems. Cell-based assays include cells naturally expressing the transporter nucleic acid or recombinant cells genetically engineered to express specific nucleic acid sequences.

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The assay for transporter nucleic acid expression can involve direct assay of nucleic acid levels, such as mRNA levels, or on collateral compounds involved in the signal pathway. Further, the expression of genes that are up- or down-regulated in response to the transporter protein signal pathway can also be assayed. In this embodiment the regulatory regions of these genes can be operably linked to a reporter gene such as luciferase.

Thus, modulators of transporter gene expression can be identified in a method wherein a cell is contacted with a candidate compound and the expression of mRNA determined. The level of expression of transporter mRNA in the presence of the candidate compound is compared to the level of expression of transporter mRNA in the absence of the candidate compound. The candidate compound can then be identified as a modulator of nucleic acid expression based on this comparison and be used, for example to treat a disorder characterized by aberrant nucleic acid expression. When expression of mRNA is statistically significantly greater in the presence of the candidate compound than in its absence, the candidate compound is identified as a stimulator of nucleic acid expression. When nucleic acid expression is statistically significantly less in the presence of the candidate compound than in its absence, the candidate compound is identified as an inhibitor of nucleic acid expression.

The invention further provides methods of treatment, with the nucleic acid as a target, using a compound identified through drug screening as a gene modulator to modulate transporter nucleic acid expression in cells and tissues that express the transporter. Experimental data as provided in Figure 1 indicates that sodium/calcium exchanger proteins of the present invention are expressed in humans in the heart, retina, kidney, fetal brain, and fetal heart. Specifically, a virtual northern blot shows expression in the fetal brain. In addition, PCR-based tissue screening panel indicates expression in brain, heart, kidney, lung, spleen, testis, leukocyte and fetal brain. Modulation includes both up-regulation (i.e. activation or agonization) or down-regulation (suppression or antagonization) or nucleic acid expression.

Alternatively, a modulator for transporter nucleic acid expression can be a small molecule or drug identified using the screening assays described herein as long as the drug or small molecule inhibits the transporter nucleic acid expression in the cells and tissues that express the protein. Experimental data as provided in Figure 1 indicates expression in humans in brain, heart, kidney, lung, spleen, testis, leukocyte and fetal brain.

The nucleic acid molecules are also useful for monitoring the effectiveness of modulating compounds on the expression or activity of the transporter gene in clinical trials or in a treatment regimen. Thus, the gene expression pattern can serve as a barometer for the continuing

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effectiveness of treatment with the compound, particularly with compounds to which a patient can develop resistance. The gene expression pattern can also serve as a marker indicative of a physiological response of the affected cells to the compound. Accordingly, such monitoring would allow either increased administration of the compound or the administration of alternative compounds to which the patient has not become resistant. Similarly, if the level of nucleic acid expression falls below a desirable level, administration of the compound could be commensurately decreased.

The nucleic acid molecules are also useful in diagnostic assays for qualitative changes in transporter nucleic acid expression, and particularly in qualitative changes that lead to pathology. The nucleic acid molecules can be used to detect mutations in transporter genes and gene expression products such as mRNA. The nucleic acid molecules can be used as hybridization probes to detect naturally occurring genetic mutations in the transporter gene and thereby to determine whether a subject with the mutation is at risk for a disorder caused by the mutation. Mutations include deletion, addition, or substitution of one or more nucleotides in the gene, chromosomal rearrangement, such as inversion or transposition, modification of genomic DNA, such as aberrant methylation patterns or changes in gene copy number, such as amplification. Detection of a mutated form of the transporter gene associated with a dysfunction provides a diagnostic tool for an active disease or susceptibility to disease when the disease results from overexpression, underexpression, or altered expression of a transporter protein.

Individuals carrying mutations in the transporter gene can be detected at the nucleic acid level by a variety of techniques. Figure 3 provides information on SNPs that have been identified in a gene encoding the transporter protein of the present invention. 140 SNP variants were found, including 6 indels (indicated by a "-") and 1 SNPs in exons. The others were found in in introns and regions 5' and 3' of the ORF. Such SNPs in introns and outside the ORF may affect control/regulatory elements. As indicated by the data presented in Figure 3, the map position was determined to be on chromosome 14 by ePCR. Genomic DNA can be analyzed directly or can be amplified by using PCR prior to analysis. RNA or cDNA can be used in the same way. In some uses, detection of the mutation involves the use of a probe/primer in a polymerase chain reaction (PCR) (see, e.g. U.S. Patent Nos. 4,683,195 and 4,683,202), such as anchor PCR or RACE PCR, or, alternatively, in a ligation chain reaction (LCR) (see, e.g., Landegran et al., Science 241:1077-1080 (1988); and Nakazawa et al., PNAS 91:360-364 (1994)), the latter of which can be particularly useful for detecting point mutations in the gene (see Abravaya et al., Nucleic Acids Res. 23:675-682 (1995)). This method can include the steps of collecting a sample of cells from a patient, isolating

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nucleic acid (e.g., genomic, mRNA or both) from the cells of the sample, contacting the nucleic acid sample with one or more primers which specifically hybridize to a gene under conditions such that hybridization and amplification of the gene (if present) occurs, and detecting the presence or absence of an amplification product, or detecting the size of the amplification product and comparing the length to a control sample. Deletions and insertions can be detected by a change in size of the amplified product compared to the normal genotype. Point mutations can be identified by hybridizing amplified DNA to normal RNA or antisense DNA sequences.

Alternatively, mutations in a transporter gene can be directly identified, for example, by alterations in restriction enzyme digestion patterns determined by gel electrophoresis.

Further, sequence-specific ribozymes (U.S. Patent No. 5,498,531) can be used to score for the presence of specific mutations by development or loss of a ribozyme cleavage site. Perfectly matched sequences can be distinguished from mismatched sequences by nuclease cleavage digestion assays or by differences in melting temperature.

Sequence changes at specific locations can also be assessed by nuclease protection assays such as RNase and S1 protection or the chemical cleavage method. Furthermore, sequence differences between a mutant transporter gene and a wild-type gene can be determined by direct DNA sequencing. A variety of automated sequencing procedures can be utilized when performing the diagnostic assays (Naeve, C.W., (1995) *Biotechniques 19*:448), including sequencing by mass spectrometry (see, e.g., PCT International Publication No. WO 94/16101; Cohen *et al.*, *Adv. Chromatogr. 36*:127-162 (1996); and Griffin *et al.*, *Appl. Biochem. Biotechnol. 38*:147-159 (1993)).

Other methods for detecting mutations in the gene include methods in which protection from cleavage agents is used to detect mismatched bases in RNA/RNA or RNA/DNA duplexes (Myers et al., Science 230:1242 (1985)); Cotton et al., PNAS 85:4397 (1988); Saleeba et al., Meth. Enzymol. 217:286-295 (1992)), electrophoretic mobility of mutant and wild type nucleic acid is compared (Orita et al., PNAS 86:2766 (1989); Cotton et al., Mutat. Res. 285:125-144 (1993); and Hayashi et al., Genet. Anal. Tech. Appl. 9:73-79 (1992)), and movement of mutant or wild-type fragments in polyacrylamide gels containing a gradient of denaturant is assayed using denaturing gradient gel electrophoresis (Myers et al., Nature 313:495 (1985)). Examples of other techniques for detecting point mutations include selective oligonucleotide hybridization, selective amplification, and selective primer extension.

The nucleic acid molecules are also useful for testing an individual for a genotype that while not necessarily causing the disease, nevertheless affects the treatment modality. Thus, the nucleic acid molecules can be used to study the relationship between an individual's genotype and the

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individual's response to a compound used for treatment (pharmacogenomic relationship). Accordingly, the nucleic acid molecules described herein can be used to assess the mutation content of the transporter gene in an individual in order to select an appropriate compound or dosage regimen for treatment. Figure 3 provides information on SNPs that have been identified in a gene encoding the transporter protein of the present invention. 140 SNP variants were found, including 6 indels (indicated by a "-") and 1 SNPs in exons. The others were found in in introns and regions 5' and 3' of the ORF. Such SNPs in introns and outside the ORF may affect control/regulatory elements.

Thus nucleic acid molecules displaying genetic variations that affect treatment provide a diagnostic target that can be used to tailor treatment in an individual. Accordingly, the production of recombinant cells and animals containing these polymorphisms allow effective clinical design of treatment compounds and dosage regimens.

The nucleic acid molecules are thus useful as antisense constructs to control transporter gene expression in cells, tissues, and organisms. A DNA antisense nucleic acid molecule is designed to be complementary to a region of the gene involved in transcription, preventing transcription and hence production of transporter protein. An antisense RNA or DNA nucleic acid molecule would hybridize to the mRNA and thus block translation of mRNA into transporter protein.

Alternatively, a class of antisense molecules can be used to inactivate mRNA in order to decrease expression of transporter nucleic acid. Accordingly, these molecules can treat a disorder characterized by abnormal or undesired transporter nucleic acid expression. This technique involves cleavage by means of ribozymes containing nucleotide sequences complementary to one or more regions in the mRNA that attenuate the ability of the mRNA to be translated. Possible regions include coding regions and particularly coding regions corresponding to the catalytic and other functional activities of the transporter protein, such as ligand binding.

The nucleic acid molecules also provide vectors for gene therapy in patients containing cells that are aberrant in transporter gene expression. Thus, recombinant cells, which include the patient's cells that have been engineered *ex vivo* and returned to the patient, are introduced into an individual where the cells produce the desired transporter protein to treat the individual.

The invention also encompasses kits for detecting the presence of a transporter nucleic acid in a biological sample. Experimental data as provided in Figure 1 indicates that sodium/calcium exchanger proteins of the present invention are expressed in humans in the heart, retina, kidney, fetal brain, and fetal heart. Specifically, a virtual northern blot shows expression in the fetal brain. In addition, PCR-based tissue screening panel indicates expression in brain, heart, kidney, lung,

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spleen, testis, leukocyte and fetal brain. For example, the kit can comprise reagents such as a labeled or labelable nucleic acid or agent capable of detecting transporter nucleic acid in a biological sample; means for determining the amount of transporter nucleic acid in the sample; and means for comparing the amount of transporter nucleic acid in the sample with a standard. The compound or agent can be packaged in a suitable container. The kit can further comprise instructions for using the kit to detect transporter protein mRNA or DNA.

# Nucleic Acid Arrays

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The present invention further provides nucleic acid detection kits, such as arrays or microarrays of nucleic acid molecules that are based on the sequence information provided in Figures 1 and 3 (SEQ ID NOS:1 and 3).

As used herein "Arrays" or "Microarrays" refers to an array of distinct polynucleotides or oligonucleotides synthesized on a substrate, such as paper, nylon or other type of membrane, filter, chip, glass slide, or any other suitable solid support. In one embodiment, the microarray is prepared and used according to the methods described in US Patent 5,837,832, Chee *et al.*, PCT application W095/11995 (Chee *et al.*), Lockhart, D. J. *et al.* (1996; Nat. Biotech. 14: 1675-1680) and Schena, M. *et al.* (1996; Proc. Natl. Acad. Sci. 93: 10614-10619), all of which are incorporated herein in their entirety by reference. In other embodiments, such arrays are produced by the methods described by Brown *et al.*, US Patent No. 5,807,522.

The microarray or detection kit is preferably composed of a large number of unique, single-stranded nucleic acid sequences, usually either synthetic antisense oligonucleotides or fragments of cDNAs, fixed to a solid support. The oligonucleotides are preferably about 6-60 nucleotides in length, more preferably 15-30 nucleotides in length, and most preferably about 20-25 nucleotides in length. For a certain type of microarray or detection kit, it may be preferable to use oligonucleotides that are only 7-20 nucleotides in length. The microarray or detection kit may contain oligonucleotides that cover the known 5', or 3', sequence, sequential oligonucleotides that cover the full length sequence; or unique oligonucleotides selected from particular areas along the length of the sequence. Polynucleotides used in the microarray or detection kit may be oligonucleotides that are specific to a gene or genes of interest.

In order to produce oligonucleotides to a known sequence for a microarray or detection kit, the gene(s) of interest (or an ORF identified from the contigs of the present invention) is typically examined using a computer algorithm which starts at the 5' or at the 3' end of the nucleotide sequence. Typical algorithms will then identify oligomers of defined length that are

unique to the gene, have a GC content within a range suitable for hybridization, and lack predicted secondary structure that may interfere with hybridization. In certain situations it may be appropriate to use pairs of oligonucleotides on a microarray or detection kit. The "pairs" will be identical, except for one nucleotide that preferably is located in the center of the sequence. The second oligonucleotide in the pair (mismatched by one) serves as a control. The number of oligonucleotide pairs may range from two to one million. The oligomers are synthesized at designated areas on a substrate using a light-directed chemical process. The substrate may be paper, nylon or other type of membrane, filter, chip, glass slide or any other suitable solid support.

In another aspect, an oligonucleotide may be synthesized on the surface of the substrate by using a chemical coupling procedure and an ink jet application apparatus, as described in PCT application W095/251116 (Baldeschweiler *et al.*) which is incorporated herein in its entirety by reference. In another aspect, a "gridded" array analogous to a dot (or slot) blot may be used to arrange and link cDNA fragments or oligonucleotides to the surface of a substrate using a vacuum system, thermal, UV, mechanical or chemical bonding procedures. An array, such as those described above, may be produced by hand or by using available devices (slot blot or dot blot apparatus), materials (any suitable solid support), and machines (including robotic instruments), and may contain 8, 24, 96, 384, 1536, 6144 or more oligonucleotides, or any other number between two and one million which lends itself to the efficient use of commercially available instrumentation.

In order to conduct sample analysis using a microarray or detection kit, the RNA or DNA from a biological sample is made into hybridization probes. The mRNA is isolated, and cDNA is produced and used as a template to make antisense RNA (aRNA). The aRNA is amplified in the presence of fluorescent nucleotides, and labeled probes are incubated with the microarray or detection kit so that the probe sequences hybridize to complementary oligonucleotides of the microarray or detection kit. Incubation conditions are adjusted so that hybridization occurs with precise complementary matches or with various degrees of less complementarity. After removal of nonhybridized probes, a scanner is used to determine the levels and patterns of fluorescence. The scanned images are examined to determine degree of complementarity and the relative abundance of each oligonucleotide sequence on the microarray or detection kit. The biological samples may be obtained from any bodily fluids (such as blood, urine, saliva, phlegm, gastric juices, etc.), cultured cells, biopsies, or other tissue preparations. A detection system may be used to measure the absence, presence, and amount of hybridization for all of the distinct

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sequences simultaneously. This data may be used for large-scale correlation studies on the sequences, expression patterns, mutations, variants, or polymorphisms among samples.

Using such arrays, the present invention provides methods to identify the expression of the transporter proteins/peptides of the present invention. In detail, such methods comprise incubating a test sample with one or more nucleic acid molecules and assaying for binding of the nucleic acid molecule with components within the test sample. Such assays will typically involve arrays comprising many genes, at least one of which is a gene of the present invention and or alleles of the transporter gene of the present invention. Figure 3 provides information on SNPs that have been identified in a gene encoding the transporter protein of the present invention. 140 SNP variants were found, including 6 indels (indicated by a "-") and 1 SNPs in exons. The others were found in in introns and regions 5' and 3' of the ORF. Such SNPs in introns and outside the ORF may affect control/regulatory elements.

Conditions for incubating a nucleic acid molecule with a test sample vary. Incubation conditions depend on the format employed in the assay, the detection methods employed, and the type and nature of the nucleic acid molecule used in the assay. One skilled in the art will recognize that any one of the commonly available hybridization, amplification or array assay formats can readily be adapted to employ the novel fragments of the Human genome disclosed herein. Examples of such assays can be found in Chard, T, *An Introduction to Radioimmunoassay and Related Techniques*, Elsevier Science Publishers, Amsterdam, The Netherlands (1986); Bullock, G. R. *et al.*, *Techniques in Immunocytochemistry*, Academic Press, Orlando, FL Vol. 1 (1982), Vol. 2 (1983), Vol. 3 (1985); Tijssen, P., *Practice and Theory of Enzyme Immunoassays: Laboratory Techniques in Biochemistry and Molecular Biology*, Elsevier Science Publishers, Amsterdam, The Netherlands (1985).

The test samples of the present invention include cells, protein or membrane extracts of cells. The test sample used in the above-described method will vary based on the assay format, nature of the detection method and the tissues, cells or extracts used as the sample to be assayed. Methods for preparing nucleic acid extracts or of cells are well known in the art and can be readily be adapted in order to obtain a sample that is compatible with the system utilized.

In another embodiment of the present invention, kits are provided which contain the necessary reagents to carry out the assays of the present invention.

Specifically, the invention provides a compartmentalized kit to receive, in close confinement, one or more containers which comprises: (a) a first container comprising one of the nucleic acid molecules that can bind to a fragment of the Human genome disclosed herein; and

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(b) one or more other containers comprising one or more of the following: wash reagents, reagents capable of detecting presence of a bound nucleic acid.

In detail, a compartmentalized kit includes any kit in which reagents are contained in separate containers. Such containers include small glass containers, plastic containers, strips of plastic, glass or paper, or arraying material such as silica. Such containers allows one to efficiently transfer reagents from one compartment to another compartment such that the samples and reagents are not cross-contaminated, and the agents or solutions of each container can be added in a quantitative fashion from one compartment to another. Such containers will include a container which will accept the test sample, a container which contains the nucleic acid probe, containers which contain wash reagents (such as phosphate buffered saline, Tris-buffers, etc.), and containers which contain the reagents used to detect the bound probe. One skilled in the art will readily recognize that the previously unidentified transporter gene of the present invention can be routinely identified using the sequence information disclosed herein can be readily incorporated into one of the established kit formats which are well known in the art, particularly expression arrays.

## Vectors/host cells

The invention also provides vectors containing the nucleic acid molecules described herein. The term "vector" refers to a vehicle, preferably a nucleic acid molecule, which can transport the nucleic acid molecules. When the vector is a nucleic acid molecule, the nucleic acid molecules are covalently linked to the vector nucleic acid. With this aspect of the invention, the vector includes a plasmid, single or double stranded phage, a single or double stranded RNA or DNA viral vector, or artificial chromosome, such as a BAC, PAC, YAC, OR MAC.

A vector can be maintained in the host cell as an extrachromosomal element where it replicates and produces additional copies of the nucleic acid molecules. Alternatively, the vector may integrate into the host cell genome and produce additional copies of the nucleic acid molecules when the host cell replicates.

The invention provides vectors for the maintenance (cloning vectors) or vectors for expression (expression vectors) of the nucleic acid molecules. The vectors can function in procaryotic or eukaryotic cells or in both (shuttle vectors).

Expression vectors contain cis-acting regulatory regions that are operably linked in the vector to the nucleic acid molecules such that transcription of the nucleic acid molecules is allowed in a host cell. The nucleic acid molecules can be introduced into the host cell with a separate

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nucleic acid molecule capable of affecting transcription. Thus, the second nucleic acid molecule may provide a trans-acting factor interacting with the cis-regulatory control region to allow transcription of the nucleic acid molecules from the vector. Alternatively, a trans-acting factor may be supplied by the host cell. Finally, a trans-acting factor can be produced from the vector itself. It is understood, however, that in some embodiments, transcription and/or translation of the nucleic acid molecules can occur in a cell-free system.

The regulatory sequence to which the nucleic acid molecules described herein can be operably linked include promoters for directing mRNA transcription. These include, but are not limited to, the left promoter from bacteriophage  $\lambda$ , the lac, TRP, and TAC promoters from *E. coli*, the early and late promoters from SV40, the CMV immediate early promoter, the adenovirus early and late promoters, and retrovirus long-terminal repeats.

In addition to control regions that promote transcription, expression vectors may also include regions that modulate transcription, such as repressor binding sites and enhancers. Examples include the SV40 enhancer, the cytomegalovirus immediate early enhancer, polyoma enhancer, adenovirus enhancers, and retrovirus LTR enhancers.

In addition to containing sites for transcription initiation and control, expression vectors can also contain sequences necessary for transcription termination and, in the transcribed region a ribosome binding site for translation. Other regulatory control elements for expression include initiation and termination codons as well as polyadenylation signals. The person of ordinary skill in the art would be aware of the numerous regulatory sequences that are useful in expression vectors. Such regulatory sequences are described, for example, in Sambrook et al., Molecular Cloning: A Laboratory Manual. 2nd. ed., Cold Spring Harbor Laboratory Press, Cold Spring Harbor, NY, (1989).

A variety of expression vectors can be used to express a nucleic acid molecule. Such vectors include chromosomal, episomal, and virus-derived vectors, for example vectors derived from bacterial plasmids, from bacteriophage, from yeast episomes, from yeast chromosomal elements, including yeast artificial chromosomes, from viruses such as baculoviruses, papovaviruses such as SV40, Vaccinia viruses, adenoviruses, poxviruses, pseudorabies viruses, and retroviruses. Vectors may also be derived from combinations of these sources such as those derived from plasmid and bacteriophage genetic elements, e.g. cosmids and phagemids. Appropriate cloning and expression vectors for prokaryotic and eukaryotic hosts are described in Sambrook et al., Molecular Cloning: A Laboratory Manual. 2nd. ed., Cold Spring Harbor Laboratory Press, Cold Spring Harbor, NY, (1989).

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The regulatory sequence may provide constitutive expression in one or more host cells (i.e. tissue specific) or may provide for inducible expression in one or more cell types such as by temperature, nutrient additive, or exogenous factor such as a hormone or other ligand. A variety of vectors providing for constitutive and inducible expression in prokaryotic and eukaryotic hosts are well known to those of ordinary skill in the art.

The nucleic acid molecules can be inserted into the vector nucleic acid by well-known methodology. Generally, the DNA sequence that will ultimately be expressed is joined to an expression vector by cleaving the DNA sequence and the expression vector with one or more restriction enzymes and then ligating the fragments together. Procedures for restriction enzyme digestion and ligation are well known to those of ordinary skill in the art.

The vector containing the appropriate nucleic acid molecule can be introduced into an appropriate host cell for propagation or expression using well-known techniques. Bacterial cells include, but are not limited to. *E. coli. Streptomyces*, and *Salmonella typhimurium*. Eukaryotic cells include, but are not limited to. yeast, insect cells such as *Drosophila*, animal cells such as COS and CHO cells, and plant cells.

As described herein, it may be desirable to express the peptide as a fusion protein. Accordingly, the invention provides fusion vectors that allow for the production of the peptides. Fusion vectors can increase the expression of a recombinant protein, increase the solubility of the recombinant protein, and aid in the purification of the protein by acting for example as a ligand for affinity purification. A proteolytic cleavage site may be introduced at the junction of the fusion moiety so that the desired peptide can ultimately be separated from the fusion moiety. Proteolytic enzymes include, but are not limited to, factor Xa, thrombin, and enterotransporter. Typical fusion expression vectors include pGEX (Smith et al., Gene 67:31-40 (1988)), pMAL (New England Biolabs, Beverly, MA) and pRIT5 (Pharmacia, Piscataway, NJ) which fuse glutathione Stransferase (GST), maltose E binding protein, or protein A, respectively, to the target recombinant protein. Examples of suitable inducible non-fusion E. coli expression vectors include pTrc (Amann et al., Gene 69:301-315 (1988)) and pET 11d (Studier et al., Gene Expression Technology: Methods in Enzymology 185:60-89 (1990)).

Recombinant protein expression can be maximized in host bacteria by providing a genetic background wherein the host cell has an impaired capacity to proteolytically cleave the recombinant protein. (Gottesman, S., Gene Expression Technology: Methods in Enzymology 185, Academic Press, San Diego, California (1990) 119-128). Alternatively, the sequence of the nucleic acid

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molecule of interest can be altered to provide preferential codon usage for a specific host cell, for example *E. coli*. (Wada *et al.*, *Nucleic Acids Res. 20*:2111-2118 (1992)).

The nucleic acid molecules can also be expressed by expression vectors that are operative in yeast. Examples of vectors for expression in yeast e.g., *S. cerevisiae* include pYepSec1 (Baldari, *et al.*, *EMBO J. 6*:229-234 (1987)), pMFa (Kurjan *et al.*, *Cell 30*:933-943(1982)), pJRY88 (Schultz *et al.*, *Gene 54*:113-123 (1987)), and pYES2 (Invitrogen Corporation, San Diego, CA).

The nucleic acid molecules can also be expressed in insect cells using, for example, baculovirus expression vectors. Baculovirus vectors available for expression of proteins in cultured insect cells (e.g., Sf 9 cells) include the pAc series (Smith *et al.*, *Mol. Cell Biol.* 3:2156-2165 (1983)) and the pVL series (Lucklow *et al.*, *Virology* 170:31-39 (1989)).

In certain embodiments of the invention, the nucleic acid molecules described herein are expressed in mammalian cells using mammalian expression vectors. Examples of mammalian expression vectors include pCDM8 (Seed, B. *Nature 329*:840(1987)) and pMT2PC (Kaufman *et al.*, *EMBO J. 6*:187-195 (1987)).

The expression vectors listed herein are provided by way of example only of the well-known vectors available to those of ordinary skill in the art that would be useful to express the nucleic acid molecules. The person of ordinary skill in the art would be aware of other vectors suitable for maintenance propagation or expression of the nucleic acid molecules described herein. These are found for example in Sambrook, J., Fritsh, E. F., and Maniatis, T. *Molecular Cloning: A Laboratory Manual. 2nd, ed., Cold Spring Harbor Laboratory*, Cold Spring Harbor Laboratory Press, Cold Spring Harbor, NY, 1989.

The invention also encompasses vectors in which the nucleic acid sequences described herein are cloned into the vector in reverse orientation, but operably linked to a regulatory sequence that permits transcription of antisense RNA. Thus, an antisense transcript can be produced to all, or to a portion, of the nucleic acid molecule sequences described herein, including both coding and non-coding regions. Expression of this antisense RNA is subject to each of the parameters described above in relation to expression of the sense RNA (regulatory sequences, constitutive or inducible expression, tissue-specific expression).

The invention also relates to recombinant host cells containing the vectors described herein. Host cells therefore include prokaryotic cells, lower eukaryotic cells such as yeast, other eukaryotic cells such as insect cells, and higher eukaryotic cells such as mammalian cells.

The recombinant host cells are prepared by introducing the vector constructs described herein into the cells by techniques readily available to the person of ordinary skill in the art. These

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include, but are not limited to, calcium phosphate transfection, DEAE-dextran-mediated transfection, cationic lipid-mediated transfection, electroporation, transduction, infection, lipofection, and other techniques such as those found in Sambrook, et al. (Molecular Cloning: A Laboratory Manual. 2nd, ed., Cold Spring Harbor Laboratory, Cold Spring Harbor Laboratory Press, Cold Spring Harbor, NY, 1989).

Host cells can contain more than one vector. Thus, different nucleotide sequences can be introduced on different vectors of the same cell. Similarly, the nucleic acid molecules can be introduced either alone or with other nucleic acid molecules that are not related to the nucleic acid molecules such as those providing trans-acting factors for expression vectors. When more than one vector is introduced into a cell, the vectors can be introduced independently, co-introduced or joined to the nucleic acid molecule vector.

In the case of bacteriophage and viral vectors, these can be introduced into cells as packaged or encapsulated virus by standard procedures for infection and transduction. Viral vectors can be replication-competent or replication-defective. In the case in which viral replication is defective, replication will occur in host cells providing functions that complement the defects.

Vectors generally include selectable markers that enable the selection of the subpopulation of cells that contain the recombinant vector constructs. The marker can be contained in the same vector that contains the nucleic acid molecules described herein or may be on a separate vector. Markers include tetracycline or ampicillin-resistance genes for prokaryotic host cells and dihydrofolate reductase or neomycin resistance for eukaryotic host cells. However, any marker that provides selection for a phenotypic trait will be effective.

While the mature proteins can be produced in bacteria, yeast, mammalian cells, and other cells under the control of the appropriate regulatory sequences, cell- free transcription and translation systems can also be used to produce these proteins using RNA derived from the DNA constructs described herein.

Where secretion of the peptide is desired, which is difficult to achieve with multitransmembrane domain containing proteins such as transporters, appropriate secretion signals are incorporated into the vector. The signal sequence can be endogenous to the peptides or heterologous to these peptides.

Where the peptide is not secreted into the medium, which is typically the case with transporters, the protein can be isolated from the host cell by standard disruption procedures, including freeze thaw, sonication, mechanical disruption, use of lysing agents and the like. The peptide can then be recovered and purified by well-known purification methods including

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ammonium sulfate precipitation, acid extraction, anion or cationic exchange chromatography, phosphocellulose chromatography, hydrophobic-interaction chromatography, affinity chromatography, hydroxylapatite chromatography, lectin chromatography, or high performance liquid chromatography.

It is also understood that depending upon the host cell in recombinant production of the peptides described herein, the peptides can have various glycosylation patterns, depending upon the cell, or maybe non-glycosylated as when produced in bacteria. In addition, the peptides may include an initial modified methionine in some cases as a result of a host-mediated process.

# Uses of vectors and host cells

The recombinant host cells expressing the peptides described herein have a variety of uses. First, the cells are useful for producing a transporter protein or peptide that can be further purified to produce desired amounts of transporter protein or fragments. Thus, host cells containing expression vectors are useful for peptide production.

Host cells are also useful for conducting cell-based assays involving the transporter protein or transporter protein fragments, such as those described above as well as other formats known in the art. Thus, a recombinant host cell expressing a native transporter protein is useful for assaying compounds that stimulate or inhibit transporter protein function.

Host cells are also useful for identifying transporter protein mutants in which these functions are affected. If the mutants naturally occur and give rise to a pathology, host cells containing the mutations are useful to assay compounds that have a desired effect on the mutant transporter protein (for example, stimulating or inhibiting function) which may not be indicated by their effect on the native transporter protein.

Genetically engineered host cells can be further used to produce non-human transgenic animals. A transgenic animal is preferably a mammal, for example a rodent, such as a rat or mouse, in which one or more of the cells of the animal include a transgene. A transgene is exogenous DNA that is integrated into the genome of a cell from which a transgenic animal develops and which remains in the genome of the mature animal in one or more cell types or tissues of the transgenic animal. These animals are useful for studying the function of a transporter protein and identifying and evaluating modulators of transporter protein activity. Other examples of transgenic animals include non-human primates, sheep, dogs, cows, goats, chickens, and amphibians.

A transgenic animal can be produced by introducing nucleic acid into the male pronuclei of a fertilized oocyte, e.g., by microinjection, retroviral infection, and allowing the oocyte to develop

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in a pseudopregnant female foster animal. Any of the transporter protein nucleotide sequences can be introduced as a transgene into the genome of a non-human animal, such as a mouse.

Any of the regulatory or other sequences useful in expression vectors can form part of the transgenic sequence. This includes intronic sequences and polyadenylation signals, if not already included. A tissue-specific regulatory sequence(s) can be operably linked to the transgene to direct expression of the transporter protein to particular cells.

Methods for generating transgenic animals via embryo manipulation and microinjection, particularly animals such as mice, have become conventional in the art and are described, for example, in U.S. Patent Nos. 4,736,866 and 4,870,009, both by Leder *et al.*, U.S. Patent No. 4,873.191 by Wagner *et al.* and in Hogan, B., *Manipulating the Mouse Embryo*, (Cold Spring Harbor Laboratory Press, Cold Spring Harbor, N.Y., 1986). Similar methods are used for production of other transgenic animals. A transgenic founder animal can be identified based upon the presence of the transgene in its genome and/or expression of transgenic mRNA in tissues or cells of the animals. A transgenic founder animal can then be used to breed additional animals carrying the transgene. Moreover, transgenic animals carrying a transgene can further be bred to other transgenic animals carrying other transgenes. A transgenic animal also includes animals in which the entire animal or tissues in the animal have been produced using the homologously recombinant host cells described herein.

In another embodiment, transgenic non-human animals can be produced which contain selected systems that allow for regulated expression of the transgene. One example of such a system is the *cre/loxP* recombinase system of bacteriophage P1. For a description of the *cre/loxP* recombinase system, see, e.g., Lakso *et al. PNAS 89*:6232-6236 (1992). Another example of a recombinase system is the FLP recombinase system of *S. cerevisiae* (O'Gorman *et al. Science 251*:1351-1355 (1991). If a *cre/loxP* recombinase system is used to regulate expression of the transgene, animals containing transgenes encoding both the *Cre* recombinase and a selected protein is required. Such animals can be provided through the construction of "double" transgenic animals, e.g., by mating two transgenic animals, one containing a transgene encoding a selected protein and the other containing a transgene encoding a recombinase.

Clones of the non-human transgenic animals described herein can also be produced according to the methods described in Wilmut, I. et al. Nature 385:810-813 (1997) and PCT International Publication Nos. WO 97/07668 and WO 97/07669. In brief, a cell, e.g., a somatic cell, from the transgenic animal can be isolated and induced to exit the growth cycle and enter G<sub>o</sub> phase. The quiescent cell can then be fused, e.g., through the use of electrical pulses, to an enucleated

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oocyte from an animal of the same species from which the quiescent cell is isolated. The reconstructed oocyte is then cultured such that it develops to morula or blastocyst and then transferred to pseudopregnant female foster animal. The offspring born of this female foster animal will be a clone of the animal from which the cell, e.g., the somatic cell, is isolated.

Transgenic animals containing recombinant cells that express the peptides described herein are useful to conduct the assays described herein in an *in vivo* context. Accordingly, the various physiological factors that are present *in vivo* and that could effect ligand binding, transporter protein activation, and signal transduction, may not be evident from *in vitro* cell-free or cell-based assays. Accordingly, it is useful to provide non-human transgenic animals to assay *in vivo* transporter protein function, including ligand interaction, the effect of specific mutant transporter proteins on transporter protein function and ligand interaction, and the effect of chimeric transporter proteins. It is also possible to assess the effect of null mutations, that is mutations that substantially or completely eliminate one or more transporter protein functions.

All publications and patents mentioned in the above specification are herein incorporated by reference. Various modifications and variations of the described method and system of the invention will be apparent to those skilled in the art without departing from the scope and spirit of the invention. Although the invention has been described in connection with specific preferred embodiments, it should be understood that the invention as claimed should not be unduly limited to such specific embodiments. Indeed, various modifications of the above-described modes for carrying out the invention which are obvious to those skilled in the field of molecular biology or related fields are intended to be within the scope of the following claims.

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#### **Claims**

That which is claimed is:

1. An isolated peptide consisting of an amino acid sequence selected from the group consisting of:

- (a) an amino acid sequence shown in SEQ ID NO:2;
- (b) an amino acid sequence of an allelic variant of an amino acid sequence shown in SEQ ID NO:2, wherein said allelic variant is encoded by a nucleic acid molecule that hybridizes under stringent conditions to the opposite strand of a nucleic acid molecule shown in SEQ ID NOS:1 or 3;
- (c) an amino acid sequence of an ortholog of an amino acid sequence shown in SEQ ID NO:2, wherein said ortholog is encoded by a nucleic acid molecule that hybridizes under stringent conditions to the opposite strand of a nucleic acid molecule shown in SEQ ID NOS:1 or 3; and
- (d) a fragment of an amino acid sequence shown in SEQ ID NO:2, wherein said fragment comprises at least 10 contiguous amino acids.
- 2. An isolated peptide comprising an amino acid sequence selected from the group consisting of:
  - (a) an amino acid sequence shown in SEQ ID NO:2;
- (b) an amino acid sequence of an allelic variant of an amino acid sequence shown in SEQ ID NO:2, wherein said allelic variant is encoded by a nucleic acid molecule that hybridizes under stringent conditions to the opposite strand of a nucleic acid molecule shown in SEQ ID NOS:1 or 3;
- (c) an amino acid sequence of an ortholog of an amino acid sequence shown in SEQ ID NO:2, wherein said ortholog is encoded by a nucleic acid molecule that hybridizes under stringent conditions to the opposite strand of a nucleic acid molecule shown in SEQ ID NOS:1 or 3; and
- (d) a fragment of an amino acid sequence shown in SEQ ID NO:2, wherein said fragment comprises at least 10 contiguous amino acids.

3. An isolated antibody that selectively binds to a peptide of claim 2.

4. An isolated nucleic acid molecule consisting of a nucleotide sequence selected from the group consisting of:

- (a) a nucleotide sequence that encodes an amino acid sequence shown in SEQ ID NO:2;
- (b) a nucleotide sequence that encodes of an allelic variant of an amino acid sequence shown in SEQ ID NO:2, wherein said nucleotide sequence hybridizes under stringent conditions to the opposite strand of a nucleic acid molecule shown in SEQ ID NOS:1 or 3;
- (c) a nucleotide sequence that encodes an ortholog of an amino acid sequence shown in SEQ ID NO:2, wherein said nucleotide sequence hybridizes under stringent conditions to the opposite strand of a nucleic acid molecule shown in SEQ ID NOS:1 or 3;
- (d) a nucleotide sequence that encodes a fragment of an amino acid sequence shown in SEQ ID NO:2, wherein said fragment comprises at least 10 contiguous amino acids; and
- (e) a nucleotide sequence that is the complement of a nucleotide sequence of (a)-(d).
- 5. An isolated nucleic acid molecule comprising a nucleotide sequence selected from the group consisting of:
- (a) a nucleotide sequence that encodes an amino acid sequence shown in SEQ ID NO:2;
- (b) a nucleotide sequence that encodes of an allelic variant of an amino acid sequence shown in SEQ ID NO:2, wherein said nucleotide sequence hybridizes under stringent conditions to the opposite strand of a nucleic acid molecule shown in SEQ ID NOS:1 or 3;
- (c) a nucleotide sequence that encodes an ortholog of an amino acid sequence shown in SEQ ID NO:2, wherein said nucleotide sequence hybridizes under stringent conditions to the opposite strand of a nucleic acid molecule shown in SEQ ID NOS:1 or 3;
- (d) a nucleotide sequence that encodes a fragment of an amino acid sequence shown in SEQ ID NO:2, wherein said fragment comprises at least 10 contiguous amino acids; and
- (e) a nucleotide sequence that is the complement of a nucleotide sequence of (a)-(d).

- 6. A gene chip comprising a nucleic acid molecule of claim 5.
- 7. A transgenic non-human animal comprising a nucleic acid molecule of claim 5.
- 8. A nucleic acid vector comprising a nucleic acid molecule of claim 5.
- 9. A host cell containing the vector of claim 8.
- 10. A method for producing any of the peptides of claim 1 comprising introducing a nucleotide sequence encoding any of the amino acid sequences in (a)-(d) into a host cell, and culturing the host cell under conditions in which the peptides are expressed from the nucleotide sequence.
- 11. A method for producing any of the peptides of claim 2 comprising introducing a nucleotide sequence encoding any of the amino acid sequences in (a)-(d) into a host cell, and culturing the host cell under conditions in which the peptides are expressed from the nucleotide sequence.
- 12. A method for detecting the presence of any of the peptides of claim 2 in a sample, said method comprising contacting said sample with a detection agent that specifically allows detection of the presence of the peptide in the sample and then detecting the presence of the peptide.
- 13. A method for detecting the presence of a nucleic acid molecule of claim 5 in a sample, said method comprising contacting the sample with an oligonucleotide that hybridizes to said nucleic acid molecule under stringent conditions and determining whether the oligonucleotide binds to said nucleic acid molecule in the sample.
- 14. A method for identifying a modulator of a peptide of claim 2, said method comprising contacting said peptide with an agent and determining if said agent has modulated the function or activity of said peptide.

15. The method of claim 14, wherein said agent is administered to a host cell comprising an expression vector that expresses said peptide.

- 16. A method for identifying an agent that binds to any of the peptides of claim 2, said method comprising contacting the peptide with an agent and assaying the contacted mixture to determine whether a complex is formed with the agent bound to the peptide.
- 17. A pharmaceutical composition comprising an agent identified by the method of claim 16 and a pharmaceutically acceptable carrier therefor.
- 18. A method for treating a disease or condition mediated by a human transporter protein, said method comprising administering to a patient a pharmaceutically effective amount of an agent identified by the method of claim 16.
- 19. A method for identifying a modulator of the expression of a peptide of claim 2, said method comprising contacting a cell expressing said peptide with an agent, and determining if said agent has modulated the expression of said peptide.
- 20. An isolated human transporter peptide having an amino acid sequence that shares at least 70% homology with an amino acid sequence shown in SEQ ID NO:2.
- 21. A peptide according to claim 20 that shares at least 90 percent homology with an amino acid sequence shown in SEQ ID NO:2.
- 22. An isolated nucleic acid molecule encoding a human transporter peptide, said nucleic acid molecule sharing at least 80 percent homology with a nucleic acid molecule shown in SEQ ID NOS:1 or 3.
- 23. A nucleic acid molecule according to claim 22 that shares at least 90 percent homology with a nucleic acid molecule shown in SEQ ID NOS:1 or 3.

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  51 CCATTTTGGG CTGGTTACCT TTGTGCTCTT CCTGAATGGT CTTCGAGCAG
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 351 TAAGAAACCC AATGGAGAAA CCAGCACAAC CACTATTCGG GTCTGGAATG
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Start Codon: 10
Stop Codon: 2773
3'UTR: 2776

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dbEST:		
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EXPRESSION INFORMATION FOR MODULATORY USE: gi 11600765 Pooled (Brain, Heart, Kidney, Lung, Spleen, Testis, I gi 318815 Fetal brain	eukocyt	:e)

<u>Tissue expression:</u>
Pooled tissues (Brain, Heart, Kidney, Lung, Spleen, Testis, Leukocyte)

```
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  151 SLIEVCGHGF IAGDLGPSTI VGSAAFNMFI IIGICVYVIP DGETRKIKHL
  201 RVFFITAAWS IFAYIWLYMI LAVFSPGVVQ VWEGLLTLFF FPVCVLLAWV
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N-glycosylation s:te
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            817-820 NUTO
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CAMP- and cGMP-dependent protein timese phosphorylation site
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            277-280 TEGD
            312-315 SRRE
            382-385 SMSE
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            460-463 TQKE
            522-525 TILD
      10
            583-586 TYGE
      11
            637-640 TMEE
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            672-675 TTVD
      13
            691-694 SWRD
      14
            713-716 SGEE
            720-723 SCFD
     16
            794-797 SVPD
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[6] PDOC00008 PS00008 MYRISTYL
N-myristoylation site
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           497-502 GMPPAI
           557-562 GARGTV
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              End
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913

2.138 Certain

>CRA|18000005047237 /altid=gi|2498054 /def=sp|P70549|NAC3 RAT SODIUM/CALCIUM EXCHANGER 3 PRECURSOR (NA+/CA2+-EXCHANGE PROTEIN 3) /org=NA+/CA2+-EXCHANGE PROTEIN 3 /dataset=nraa /length=927 Length = 927 Score = 1828 bits (4682), Expect = 0.0Identities = 897/927 (96%), Positives = 911/927 (97%), Gaps = 6/927 (0%) Frame = +1MAWLRLQPLTSAFLHFGLVTFVLFLNGLRAEAGGSGDVPSTGQNNESCSGSSDCKEGVIL 189 Query: 10 MAWLRLQPLTSAFLHFGLVTFVLFLNGLRAEAG DVPS GQNNESCSGSSDCKEGVIL MAWLRLQPLTSAFLHFGLVTFVLFLNGLRAEAGDLRDVPSAGQNNESCSGSSDCKEGVIL 60 Sbjct: 1 Query: 190 PIWYPENPSLGDKIARVIVYFVALIYMFLGVSIIADRFMASIEVITSQEREVTIKKPNGE 369 PIWYPENPSLGDKIARVIVYFVALIYMFLGVSIIADRFMASIEVITSQEREVTIKKPNGE PIWYPENPSLGDKIARVIVYFVALIYMFLGVSIIADRFMASIEVITSQEREVTIKKPNGE 120 Sbjct: 61 TSTTTIRVWNETVSNLTLMALGSSAPEILLSLIEVCGHGFIAGDLGPSTIVGSAAFNMFI 549 Query: 370 TSTTTIRVWNETVSNLTLMALGSSAPEILLSLIEVĆGHGFIAGDLGPSTIVGSAAFNMFI Sbjct: 121 TSTTTIRVWNETVSNLTLMALGSSAPEILLSLIEVCGHGFIAGDLGPSTIVGSAAFNMFI 180 Query: 550 IIGICVYVIPDGETRKIKHLRVFFITAAWSIFAYIWLYMILAVFSPGVVQVWEGLLTLFF 729 IIGICVYVIPDGETRKIKHLRVFF+TAAWS+FAYIWLYMILAVFSPGVVQVWEGLLTLFF Sbjct: 181 IIGICVYVIPDGETRKIKHLRVFFVTAAWSVFAYIWLYMILAVFSPGVVQVWEGLLTLFF 240 FPVCVLLAWVADKRLLFYKYMHKKYRTDKHRGIIIETEGDHPKGIEMDGKMMNSHFLDGN 909 Query: 730 FPVCVLLAWVADKRLLFYKYMHK+YRTDKHRGIIIETEG+HPKGIEMDGKMMNSHFLDGN Sbjct: 241 FPVCVLLAWVADKRLLFYKYMHKRYRTDKHRGIIIETEGEHPKGIEMDGKMMNSHFLDGN 300 Query: 910 LVPLEGKEVDESRREMIRILKDLKQKHPEKDLDQLVEMANYYALSHOOKSRAFYRIOATR 1089 L+PLEGKEVDESRREMIRILKDLKQKHPEKDLDQLVEMANYYALSHQQKSRAFYRIQATR Sbjct: 301 LIPLEGKEVDESRREMIRILKDLKQKHPEKDLDQLVEMANYYALSHQQKSRAFYRIQATR 360 Query: 1090 MMTGAGNILKKHAAEQAKKASSMSEVHTDEPEDFISKVFFDPCSYQCLENCGAVLLTVVR 1269 MMTGAGNILKKHAAEQAKK +SMSEVHTDEPEDF SKVFFDPCSYQCLENCGAVLLTVVR Sbjct: 361 MMTGAGNILKKHAAEQAKKTASMSEVHTDEPEDFASKVFFDPCSYQCLENCGAVLLTVVR 420 Query: 1270 KGGDMSKTMYVDYKTEDGSANAGADYEFTEGTVVLKPGETQKEFSVGIIDDDIFEEDEHF 1449 KGGD+SKTMYVDYKTEDGSANAGADYEFTEGTVVLKPGETQKEFSVGIIDDDIFEEDEHF Sbjct: 421 KGGDISKTMYVDYKTEDGSANAGADYEFTEGTVVLKPGETQKEFSVGIIDDDIFEEDEHF 480 Query: 1450 FVRLSNVRIEEEQPEEGMPPAIFNSLPLPRAVLASPCVATVTILDDDHAGIFTFECDTIH 1629 FVRLSNVR+EEEQ EEGM PAI NSLPLPRAVLASPCVATVTILDDDHAGIFTFECDTIH Sbjct: 481 FVRLSNVRVEEEQLEEGMTPAILNSLPLPRAVLASPCVATVTILDDDHAGIFTFECDTIH 540 Query: 1630 VSESIGVMEVKVLRTSGARGTVIVPFRTVEGTAKGGGEDFEDTYGELEFKNDETVKTIRV 1809 VSESIGVMEVKVLRTSGARGTVIVPFRTVEGTAKGGGEDFEDTYGELEFKNDETVKTIRV Sbjct: 541 VSESIGVMEVKVLRTSGARGTVIVPFRTVEGTAKGGGEDFEDTYGELEFKNDETVKTIRV 600 Query: 1810 KIVDEEEYERQENFFIALGEPKWMERGIS-----DVTDRKLTMEEEEAKRIAEMGKPVL 1971 KIVDEEEYERQENFFIALGEPKWMERGIS +VTDRKLTMEEEEAKRIAEMGKPVL Sbjct: 601 KIVDEEEYERQENFFIALGEPKWMERGISALLLSPEVTDRKLTMEEEEAKRIAEMGKPVL 660 Query: 1972 GEHPKLEVIIEESYEFKTTVDKLIKKTNLALVVGTHSWRDQFMEAITVSAAGDEDEDESG 2151 GEHPKLEVIIEESYEFK+TVDKLIKKTNLALVVGTHSWRDQFMEAITVSAAGDE+EDESG GEHPKLEVIIEESYEFKSTVDKLIKKTNLALVVGTHSWRDQFMEAITVSAAGDEEEDESG 720 Query: 2152 EERLPSCFDYVMHFLTVFWKVLFACVPPTEYCHGWACFAVSILIIGMLTAIIGDLASHFG 2331 EERLPSCFDYVMHFLTVFWKVLFACVPPTEYCHGWACF VSILIIGMLTAIIGDLASHFG Sbjct: 721 EERLPSCFDYVMHFLTVFWKVLFACVPPTEYCHGWACFVVSILIIGMLTAIIGDLASHFG 780 Query: 2332 CTIGLKDSVTAVVFVAFGTSVPDTFASKAAALQDVYADASIGNVTGSNAVNVFLGIGLAW 2511 CTIGLKDSVTAVVFVAFGTSVPDTFASKAAALQDVYADASIGNVTGSNAVNVFLGIGLAW Sbjct: 781 CTIGLKDSVTAVVFVAFGTSVPDTFASKAAALQDVYADASIGNVTGSNAVNVFLGIGLAW 840 Query: 2512 SVAAIYWALQGQEFHVSAGTLAFSVTLFTIFAFVCISVLLYRRRPHLGGELGGPRGCKLA 2691 SVAAIYWA+QGQEFHVSAGTLAFSVTLFTIFAFVC+SVLLYRRRPHLGGELGGPRGCKLA Sbjct: 841 SVAAIYWAMQGQEFHVSAGTLAFSVTLFTIFAFVCLSVLLYRRRPHLGGELGGPRGCKLA 900 Query: 2692 TTWLFVSLWLLYILFATLEAYCYIKGF 2772 TTWLFVSLWLLY+LFATLEAYCYIKGF Sbjct: 901 TTWLFVSLWLLYVLFATLEAYCYIKGF 927 (SEQ ID NO:4)

Hmmer search results (Pfam):

BLAST Alignment to Top Hit:

FIGURE 2, page 3 of 4

Scores fo	or sequer Descript	nce far tion	nily cl	lass	sifica	tion (	core	includ	es all do Score	mains): E-value	N
PF01699 PF00324 PF01971	Sodium/o Amino ao Protein	cid per	mease	-	•	tein			294.6 2.8 2.7	1.2e-84 5.9 8.7	2 1 1
Parsed for Model	or domain		•		hmm-f	hmm-t		score	E-value		
PF01699 PF01971 PF00324 PF01699	1/2 1/1 1/1 2/2	118 644 851 757	257 670 877 905	• •	12 193 472 1	222	.;	121.3 2.7 2.8 181.4	1.8e-32 8.7 5.9 1.5e-50		

1 TTGGATGAGA TCTAAAGCAT TATTAAGAGT GGGGAGTGCA AAGAAGAAAC 51 CCTCATTTCA AAGATGAATG AGAATAATGG CATGTACAAA GGTCCTGGGG 101 TGGACAGTCA CTTGGTATAA TCCAAGAGTG AACCTGAAGG CTATTGTTGT 151 TGAAATGTAA TAAGGGAGAG AGTGACGGGA TGAAGGGGGA TGAGTGGGAA 201 GCAGTGAATT CCTGCAAGGC TTTGAAGGTC ATGGGAAAGA ATTTGGTCTT 251 TATATCAAGA GCAAGAGAAG ACTACTAAAG GGCTTCAAAC AGGGGAGCGA 301 TATGCTTAAG TCTGTTTGTT TGTTTTTTTA AAAAAAGATT ACGGTGGCTA 351 TATGAGGAAA GTGGAATTGA GAACTAGCGA GAGTTGGAGT GGTGAGCTCC 401 ATTAGGAGGC TACTGAAGTA GATTCATGAG GTAAGGAGTG ATGGTGGCCT 451 GGGCTGGGAT GATGGTGGTA GAAATGGAGA AAGAGTTGAT AGGATTTAGT 501 GATTGGATAA GGGACAGAAG AGAGATGAAG GCTTTCAGAC TAACATCTGC 551 TTTCTAACAT GAGTAACTGG GTGGCTGAAG ATGCTATTTT CTGAGCTGGG 601 AAACAGGAGA AAAAGGAGCA AATATGGGGG ATGAAGACTT TGAGTCTTTA 651 AGGTGCTGTA CAAACACAAA TCAGCATTCC TTTATTACTA AGGGTATCCC 701 ACACAGTTGT AGCAGAGGGA GAAAGATCGC CCCCCCCCA CTTTTTTTT 751 TTTTTTAGCT ATTCCATGGT ATTTCATTC TCATCCCACC CAAATGAGGC 801 AGTGAGTGGT AAGATGAGTA TATAATAGTT TCAATTGCAT TTCATCCCAT 851 TCTTCTGAGC TCAAGCTCAC CTTTTAGTGG TTTGAGGCCA GTAGATGAAG 901 CTGCATATCA CCCCCAAAAT CTTGTCTCTA GTTTAACAAA ACTTATTTGA 951 GAGACATTTG CATGTTTTAT TAATAATGAT TTTTACCACT TGTTCCTTTC 1001 CATGITIGGG TITGAAATTI GAGTGGCTGG CGGATGATCA TCTTCCTGTT 1051 ACTGCCTGCT TAAACTGCTC ATAAGCAGGT TTTACTGGAG GGCTCAGAGC 1101 TGCTGTGAAC TTGGTCTTGG GCACAACTTA CATGGCCTCT GTTTGGCTAT 1151 GGGCTGGGTG GCATTCACCA TTTATCAACT CTTTTGATTT CCCAAGCTAT 1201 CTCAGAATTA TAGCTTGCCT CCAGAAGTCT TGCATTCGGG GAGGAAGTTT 1251 CTTTCCAAGG GAGCTCAGTT TTCAAGGTTT ATTGCTCTGT TTAATGGATG 1301 AGATCTAAAG CATTATTAAG AGTGGGGAGT GCAAAGAAGA AACACTCATT 1351 TCAAAATCGA TTGAGAATAA TGGCATGTAC AAAGGTCCTG GGGTGGACAG 1401 TCACTTGGTA TAATCCTGGA GTGAACATGA AGGCCAAGGA AATATGTATA 1451 CATTAAACAG AGCAAGGTTT TCAATTTTCT GGGGACTAGT CCATGAAAAT 1501 TCAATTCAAT ATACTCTCTT GCAAACCTAT GTTATCCAAG ATACTCAAGT 1551 ATAATGACAA CAGGGTAAGG AAGTCCGAAC ACCCCAGAAA CAGTATAAAT 1601 GGGCATGAAG ATTCAGGTTA TACATGGCCT ATTTTAAGTT GCTTCTTGAG 1651 AACTCTCACA GGTAATACCA GTTTGGGAGA CAGGACTTGA AGGCTATTGC 1701 TGCATTTCCA TCCCCAGTAT TCCCAGCTAT TTCAAGCCAT TTTTCAACGG 1751 AGTCTCCACC AGATGGTTTG GAGGACAGAG CAGCTATTTG TGCCTCCCAT 1801 TGACATCTAT TTTTCCAAGT GAGAGACTGC CCCATATGTT AGTGCAATAT 1851 GTCACTGGAG GTGAAGCATC AGTTGTATTG GTGGGAACCT GCCGTTTGCT 1901 GTCCCCTTTT TCCTCATGCC TTTTCCTGCC TCTCTGATCT TTTCTAGGTC 1951 TCTGGCCTAT CAGGAGGACA ACTGGTGCTG CAATAGAAGC CAGTGGCTAA 2001 GTCTCGTGTA TGGCGTGGTT AAGGTTGCAG CCTCTCACCT CTGCCTTCCT 2051 CCATTTTGGG CTGGTTACCT TTGTGCTCTT CCTGAATGGT CTTCGAGCAG 2101 AGGCTGGTGG CTCAGGGGAC GTGCCAAGCA CAGGGCAGAA CAATGAGTCC 2151 TGTTCAGGGT CATCGGACTG CAAGGAGGGT GTCATCCTGC CAATCTGGTA 2201 CCCGGAGAAC CCTTCCCTTG GGGACAAGAT TGCCAGGGTC ATTGTCTATT 2251 TTGTGGCCCT GATATACATG TTCCTTGGGG TGTCCATCAT TGCTGACCGC 2301 TTCATGGCAT CTATTGAAGT CATCACCTCT CAAGAGAGGG AGGTGACAAT 2351 TAAGAAACCC AATGGAGAAA CCAGCACAAC AACTATTCGG GTCTGGAATG 2401 AAACTGTCTC CAACCTGACC CTTATGGCCC TGGGTTCCTC TGCTCCTGAG 2451 ATACTCCTCT CTTTAATTGA GGTGTGTGT CATGGGTTCA TTGCTGGTGA 2501 TCTGGGACCT TCTACCATTG TAGGGAGTGC AGCCTTCAAC ATGTTCATCA 2551 TCATTGGCAT CTGTGTCTAC GTGATCCCAG ACGGAGAGAC TCGCAAGATC 2601 AAGCATCTAC GAGTCTTCTT CATCACCGCT GCTTGGAGTA TCTTTGCCTA 2651 CATCTGGCTC TATATGATTC TGGCAGTCTT CTCCCCTGGT GTGGTCCAGG 2701 TTTGGGAAGG CCTCCTCACT CTCTTCTTCT TTCCAGTGTG TGTCCTTCTG
2751 GCCTGGGTGG CAGATAAACG ACTGCTCTTC TACAAATACA TGCACAAAAA 2801 GTACCGCACA GACAAACACC GAGGAATTAT CATAGAGACA GAGGGTGACC 2851 ACCCTAAGGG CATTGAGATG GATGGGAAAA TGATGAATTC CCATTTTCTA 2901 GATGGGAACC TGGTGCCCCT GGAAGGGAAG GAAGTGGATG AGTCCCGCAG 2951 AGAGATGATC CGGATTCTCA AGGATCTGAA GCAAAAACAC CCAGAGAAGG 3001 ACTTAGATCA GCTGGTGGAG ATGGCCAATT ACTATGCTCT TTCCCACCAA 3051 CAGAAGAGCC GCGCCTTCTA CCGTATCCAA GCCACTCGTA TGATGACTGG 3101 TGCAGGCAAT ATCCTGAAGA AACATGCAGC AGAACAAGCC AAGAAGGCCT 3151 CCAGCATGAG CGAGGTGCAC ACCGATGAGC CTGAGGACTT TATTTCCAAG 3201 GTCTTCTTTG ACCCATGTTC TTACCAGTGC CTGGAGAACT GTGGGGCTGT 3251 ACTCCTGACA GTGGTGAGGA AAGGGGGAGA CATGTCAAAG ACCATGTATG 3301 TGGACTACAA AACAGAGGAT GGTTCTGCCA ATGCAGGGGC TGACTATGAG 3351 TTCACAGAGG GCACGGTGGT TCTGAAGCCA GGAGAGACCC AGAAGGAGTT 3401 CTCCGTGGGC ATAATTGATG ACGACATTTT TGAGGAGGAT GAACACTTCT 3451 TTGTAAGGTT GAGCAATGTC CGCATAGAGG AGGAGCAGCC AGAGGAGGGG 3501 ATGCCTCCAG CAATATTCAA CAGTCTTCCC TTGCCTCGGG CTGTCCTAGC 3551 CTCCCCTTGT GTGGCCACAG TTACCATCTT GGATGATGAC CATGCAGGCA 3601 TCTTCACTTT TGAATGTGAT ACTATTCATG TCAGTGAGAG TATTGGTGTT
3651 ATGGAGGTCA AGGTTCTGCG GACATCAGGT GCCCGGGGTA CAGTCATCGT 3701 CCCCTTTAGG ACAGTAGAAG GGACAGCCAA GGGTGGCGGT GAGGACTTTG 3751 AAGACACATA TGGGGAGTTG GAATTCAAGA ATGATGAAAC TGTGTAAGTA 3801 ACCTTCCTGT ATTCTGCCCC TCCCTGACCC CATCTTTTGC CATCTCTTTC

FIGURE 3, page 1 of 57

3851	TGTCTTTCTG	TACTGCACTT	TACAACATTT	CCTTGTGTTT	GTGTTAATGT
3901	CAAACTTTGG	TTCCATCACA	GGTATGCAGG	· ATCAGCAGAC	ACCACTGGAC
3951	AGGTTCTGCT	TCCAAACTCT	TCTTCAGTTT	TCTCACTTTA	AATTGTTTCT
4001	GGGCAAGGAA	TCCTGTGACA	AGAGCTAAGG	ACACAAAACA	TTTTCTTCTC
4051	TGAAACACAA	AATGATAGCT	GGTGGAGCTG	TGGGATGACA	GAAGTTTTGT
4101	GATATCAGAT	TTTGGAGAAT	TCTTGTGACT	AAGAAGGACT	AGAGAACTGC
4151	TTGGGCCTCT	TCTTCCTCCC	TTCCTCATAT	GAAGGGTATC	TATGAGCTTT
4201	GAAACCAATC	CTTTCCATTC	TGGGCAGCAA	TAGCCCATCA	GAACATTCTA
4251	AAGAAAACAA	GTGGCATTGG	CTTTGTTCCC	TGGTACTATA	TTGCCAGTCT
4301	CACTGTGTAA	CCAGATTCCA	GGCACGTCTT	CTTTAATTTG	GAAATTGCAA
4351	AATTGATAGA	AATTTAGCAA	TCTTTTTAAA	TGACCATAGA	CTATTTAATG
4401	GTGTGAGGCT	TGCCCAGCCT	AGTTGAATTG	AGTCAGTATG	GTTTGGATAC
4451	TGGAAAGTAT	CTTGGAGAAG	CAGAGCTCCC	AGGGCAGTGG	CTACTTGTCT
4501	TTAGTCACAG	GTCTAAGCTC	CAAAATCTGG	TGAAGCAGTG	AAGGAGAAAC
4551	ATCCTAGGAA	TTGTGGGAGG	AAATATATCT	TCTGTGTGGT	CCTCTCTTTT
4601	CACAGTCTAG	GACTCTCCTG	AAGTACCTCT	TCTTGGGCTA	CTGCCCCATT
4651	CAGCCCTTCA	GAAACTGTGG	GTATTACACT	TCTGTCACCT	CTATTACCCT
4701	AAGGCCTCTG	CCCATTGAAC	CCTCTTGCAA	ATTGGTTATT	CTGTCCTTTT
4751	TCCAGTTGGA	TAGCTTTAAA	AGGGAAAGCA	GAATGACTTT	CCTCAGGATT
4801	TGTAGCTTAT	GAGAAAGTAG	ACTTTCTTGG	GTGGCCTAGA	AGGTTGGAGA
4851	AGACAAACGG	GAACTTCCTC	TGAATGACTG	AACATATCCA	CAAATAATAA
4901	GCGTGGCAGG	AGATGGTGTG	AAGAGTAAAA	GGAGCATATA	GGAAGTTGTG
4951	TGTGTGGGGT	GTCTGTTTCA	AGAACCTGCT	AATTATACCT	TCAGTAAGAA
5001	ATGAAGCCAT	ACAACCTCTA	GAAGAGGAGG	AGGAAGGAAC	TCATGGAAAA
5051	GTGGGGAGCC	ATAGAAGCTA	GGGAGAGGTG	TCCTAGGAGT	GCTTCTGCCC
5101	AGGTCCAGCC	ATGAGACAGA	GCTCAAAAAG	AGCTGGGCAC	TGCTGGTGAC
5151	AGAACTGAGT	GACCCGGGGG	ATCCTGCATC	TGTTCTTACT	CAATCCCTTC
5201	TTAATAATGT	GACTTGGGGC	AGGTCATTTA	TTGGTTCTGG	<b>AACTTAACTT</b>
5251	TCTGATATGC	AAACTGGGAA	TAACAATACT	TTCCTTGCCT	GGAGGCAAGG
5301	TCAGTCCTTT	TTGCAGTTCC	TTCCAGCTCT	AAGATTTTCT	GAACCATAGA
5351	CATAAGCACT	CAGTGTAGGT	CATATTCGCA	CTTGCCAAAA	<b>ATGGATCAGG</b>
5401	GAATATTGTC	TCCTGAAGGG	AAATGGCCAT	TGACAAATTG	<b>ATTTATTAGA</b>
5451	GCTCTGTTTA	GTCATTTTGC	TGGGAAGGAT	AATCATTTGT	TAACGTAAGT
5501	AGAAACCTGT	GCCTTCTGGA	GAATACTATC	CATTTATATG	TACTCTGGGG
5551	AGAGTGTTTA	TACATACAAA	TGAAGGACAG	GGCTTCACTG	<b>GGAAAACAAA</b>
5601	CTCCATGGAA	TTTCACATGA	TTATCGCGAT	GTCAGTGTGG	AAGAAGATAT
5651	GGTAAGGCAT	TAAATGACAT	TAAGACCACA	AAATTTGCCA	TAATTTGACG
5/01	GACTTGTGGT	TCTTCTGATT	CAGAACCCTT	TCTACCCATG	TCACGGATAG
2/27	GTAGTTTTTC	AGAGATCAGA	GGCTTAGTTC	ATTCTATTAA	TTTCCTCATT
2801	CTATTAATAA	TCAATTATGC	ACCTAGGGTC	TCTGAATACG	ACTAAACCTT
5001	CCTCAAACTT	ATTTGCATTT	TCAGTTTGTA	TAATATCTTG	GTGCAAATGA
5901	GCCTCGCAAA	TGATCACTTC	TGGGTAATAC	TCATTCTAAA	GGTATGTCAA
5001	CCTTGAGAAT	TCTGGTCTAG	ATATTCTAGG	GTTTGGTGAA	CAAATCTATG
6051	TTCCCATCCA	AAAAAAAA	TITATTTTT	AGACTTCATT	CATTGCAGAA
6101	TAATGAGTCC	AMAACCTGCT	CATCTGTTCT	CACGTGGCAC	CCCTATTCTT
6151	GATATTTTAA	ATTGCAATTT	TACAACTAGA	GGCAGTATTA	CGGAGCAGAA
6201	AAATCGTGGG	TACTARGIAC	TCTGGGTTAG	GATTCTGGCT	CCACTACTGA
6251	TTTAATAATG	ATCCCCATA	AAATTTTATT	AACCTATGAA	ATTATTTCCT
6301	CATTGGCAAA	TTCTNTCCCC	TAATATCTCT	CTTGCAGGGC	CATTATGACG
6351	ATTCAAGGTA	TCATACGAAG	TGTACCTGGT	ACACGGTATA	TGCTCAGGAA
6401	ACAAGACTCT CACTGTGGAG	TOATAGTAAT	ATTGACGAAT		
					CCTAAGTAAT
6501	GAGCATGCCA AATATTAGAG	CTACAAACCA	ACTATGAAGA	GTACTTACCT	AAACTCATAA
6551	TAAATTTTAA	TGTGTTGCTC	AATCATCCAC	CCAMCTOCAG	TCATGGTTCT
6601	TTAGGATCCA	GGAGGTCTAG	CCCACCCAMM	CACAMMOOCC	GAAGGGCAGA
6651	AGTTCTGGAT	GCTGCGGGCC	CCNACTORCA	CCTCARACCT	MORCANGO
6701	TTGACCAAAC	CAGGAGACCC	ACCANACANC	TCCTTTTTTTC	CACAAGCTC
6751	TTAATTGAAT	AATGATTGTT	TCCTCTTTA	TOGITITICA	CARROCCAAR
6801	TTAGCAAGAA	CCAGAGGCTG	TGCTCTTTAA	CACACCACTC	TCCAATGCCAAT
6851	AATGGATAGC	TTCAGGGTAC	TTGGACAAAG	TTCCAACAGIC	TCCMMMCMAA
6901	TCTCTCCCTC	TTTGTATAGC	TTTTTTTTCCC	TACCAACCCT	CCMACTAGE
6951	AAAATCTGCC	CTCACTATAC	TCCCCTAAAT	ATTA ATTC A A CTT	MCD CCCCD CC
7001	CCTGTGCTCT	ATCAATAATA	TAGGATCCAC	CAATTCAAGI	COMMOCOMMO
7051	ATGCTTTACT	TCTTCAAAGG	TECTTTTA	ACCATCCAAC	DAMCCARARC
7101	CACGAGCTTT	GGDATATCAA	ACCACATCTC	AGCATGGAAG	MATGGAAAAG
7151	CTTTTAACAA	GTCACATCAC	TTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTT	ACCACCATOAC	TETTECACACA
7201	CAGAAATAAT	ATTCTCTATC	CTTCAACCCA	ATACTARATOR	TANCONCAC
7251	AAAAATGCAC	AGTGCCTTCT	CCTACATCCT	CTTCIAAATA	TCAACAAACA
7301	TTTGTTAGAT	ATTTGCTATC	TACTACCTAC	ATTACTOAT CAT	ACTOCCOOTE
7351	AATAAGTGAA	TAAGACAAGC	TCACAGCIAC	CCCCTCX ACC	ACTOGGGTTA
7401	CAAGTGGAGA	GGATCAAAGC	GTACACITOA	DTCDACCARC	CTCL LACTGT
7451	TGGTATGGCT	GAGATGGATT	GAATAAACCA	CCDDTCTCTC	CTCCCTCC
7501	TGTGTGTGT	ACCACTGAGG	ATTCTADATT	AACCTTCATT	DICCOLOCAA
7551	TAGTGACAGA	GGTGAAGTGG	GGATAGGTAC	ATCATTANTO	TACATCCATA
7601	TTACAATGAA	ACCTTAACAT	TTAAGAGGGA	TATTATTATT	THOMICONIA
7651	TCCAGAAGAA	TCCTCACCTT	TGCAACCATC	ACTATACTON'	CTTTCHIGH
					CIICIIGAGA
		Т			^

FIGURE 3, page 2 of 57

7701	ATTATGGCCT	TTAAGACTGT	AGCATGCAAT	GACAAAACCT	CACAGAGGTA
7751	TGGGTTCTGC	CCGCACACTA	ATTTCACTCA	TTAAACAAGT	GACTGGCTCC
7801	TATATCCCAG	GCTCTCAGCA	CGCCTTTGCA	AAATAACAGA	TTATTGCAGC '
7851	TCTTGGACCT	TTGATGCCTC	TGGGAATAGT	CAAAGCCACA	GATGTCAAAT
7901	ATGTAAATGC	CAAGATCTAT	TATAATTAAA	TAGTGCAGGC	CTCCTTCAAA
7951	GAAAAAAAGC	ATGTTGGCTG	TGCTGCACGT	TCTCCAACCA	AATCAGAATG
		AAGGTATCTG			
		GGAAGGTAAC			
		GTTGGAAAGG			
		TTATTCAATG			
. –		TTTTATCCAT			
		GGTAAAAGAT			
		TCTTAACTAA			
		CACCTTGAAC			
		GTCTTTCTCT			
8451	TTGGAACATC	TTTTAGGCAG	TGCTGGTGAA	CTTCAGGCTA	GGCCTTGTTC
8501	CATGAAATAA	TAAAAATTTT	CAAAATAATG	CAGACCATTC	CCTTCCAGGG
8551	ATGCTTTCTC	TGTAATGTTT	TAACCCCAAG	AAATCTTTCT	GTAAAAATCT
8601	ATAAAAATCT	GGAGTGTTCC	AGGATACAAT	TTGCACATTC	TCCAATTTAA
8651	CTAAAACACA	ATCGATTTTT	TGTTTTCTTT	TTCTTTGGCT	TAGCAAGGTT
8701	TTAAGATAGT	CTCTTTCTGG	CCACAGAGGG	AGATGATTTG	CCTCTAGAAT
		TGCTTGAGAG			
		AGAGGTGGTG			
		TTCTGCCTTC		•	
		GTCAGCCACT			
		TTGGCACAGA			
		GAATGGGGAT			
		ACCACAATTG			
		GACTTTTTT			
		TGTCGCCCAG			
		GCCTCCCGGG			
9251	AGTAGCTAGG	CCTAATATAT	ATATATTATA	CATATATATT	TATATTTATA
9301	TATATATATA	CCACCACGTC	CGGCTAATAT	ATATTTATAC	TTTTTTTTT
9351	TAGTAGGAAA	GGGGTTTCAC	CATGTTAGCC	AGTATGGTCT	CGATCTCCTG
9401	ACCTCGTGAT	CCACCAGCCT	CAGCCTCCCA	AAGTGCTGGG	ATTACAGGCG
9451	TGAGCCACCG	TGCCCGACCA	TGCTATGTAA	ACTTTTTAGC	AGAAGCTTTA
9501	GCTATTGTGT	CCCGAAGGGC	CCCAGGTCAT	GATGAAATGT	CTTTTTTTT
9551	TTTTGTCTCT	TTTCTTCTTA	ATTACTGAGA	CTGTCAAAGA	ATATGTCAAA
		TTCCAACTCC			
		ACATCTGCAA			
		TTAGGGACTA			
		CCTACCTAAG			
		GCCCTGGTCT			
		ATCCACAAGA			
		AGATGCTCAA			
		AGGCATATTC			
		ATCTCATTCT			
		CCAGGTTCAA			
		AGACGCATGT			
		AGTTTCACCA			
10201	CTCAAGTGAT	CCGCCCACCT	TGGCCTCCCA	AAAAGCTGGG	ATTACAGGCG
10251	TGAGCTACCA	CGTCCAGCCC	CCCATATACT	TTAAATCATC	TCTAGATTAC
10301	TTATAATACC	TAATACAATG	TAAATGTTAT	ATAGTTGTTT	TAATGTATTG
10351	CTTTTTTTAT	TTGTATTGTT	TTTTATTGCT	GTATTATCCT	TTTTTATGTT
10401	TTATTTTTTC	AAATATTTTC	TACCCGTGGC	ACCCACAGTT	GGTTGGTGGA
10451	ACCTGCGGTT	GGTGGAGCCC	ATGGATGTGA	AGGGCTGATA	GTATGAGAAA
10501	ACTCAGAGGT	GCAGAGTTGG	AGAGCACATC	GGGGAGAATG	TCAGCATGGG
		CACACTGTGG			
		GGTCTCATCC			
		ATTGGAGCTT			
		ATGGGCGTAG			
		CCTCATCCAC			
		AATGAGGAGG			
		GTTTTTGGAT			
		CCCTGTCAAC			
		ACTTATTCAA			
11001	TTTTTTGTTT	TTTTTTTTT	GACAGAGTCT	TGCTCTGTTG	CCCAGGCTGG
11051	AGTGAAGTGA	AGTGGCATAA	TCTGAGCTCA	CTGCAACCTC	TGCTTTCGAG
11101	TTCAAGCGAT	TCTCATGCCT	CAGCCTCCTG	CATAGCTGGG	ACTACAGGCA
		TGCCTGGCTA			
		GCCAGGCTGA			
		TCCCAAAGTG			
		TGTTGGTTTT			
		GAAAGTTTGA			
		AGAAAGAGAG			
		TCTGCACATT			
11201	1 GAGGATTTT	GCTAAATAAC	CHIGGAGGAA	AGCACTAGAC	AAATATTTTC
		7		T 4	

FIGURE 3, page 3 of 57

11551	AGATGGCATG	AGAGTTATCA	TTCATAGGAA	TTATATTTCC	ACTCCTACCA
11601	CTTACTGGGG	ACCCAAGTAA	GAAATTACTT	GGATAAGCAG	ACCACAATTT
11651	AAAGTTGAAT	GTGGTGGAAC	TTATTATGGA	AAAAATATGT	TTTTCTGAAA
11751	ACTGGATATG TACTCTTCCT	TGTATATATA	TAAGTTCAGT	TGTCATTTTG	GAACCATCCT
	TGCCTGGACC	CAATTCAGTT	ACCTTTTCCT	GGGTACCCTT	TGACTAACTC
11851	CAGTTATTTG	TGGAGTGTAT	AGAAACCACT	CTATTGTAGG	TTCTTTACTT
11901		AAATAAGTGA	CATCCAAATA	GTAACTTAAT	ATTCCAAATA
11951	TGGCTGCAAA	ACAAATTGTC	GATTATGGAT	GACTACTACT	GCCATCTCTC
12001	CATACCAGTC	CATCTTCTGC	CAGGCTGTTT	GGTCTTGATT	TGTCGACCTT
12051 12101	TOTCCAAACG	CCCCATGTAT	TCCACATGAC	CTTCACCAAC	CCCACTTCTA
12151		TAACCGAGCC	TTGTGGGGAT CTGGCTGCGG	GCAGATGTAT	TCTGCCACCA
	TATTTCCATT	CTTACACCCT	ACTTCATGTT	TGTACACTAT	TTGTTCACAT
12251	TTGCTGTCTC	TTCTAAACAT	TCTTTGCTGC	ATCCACTTTT	TCTCTATTTG
12301	TGCTCTAGGT	GCTGCAGAGG	CTAATGCTGG	GTTTCCTTTC	ATTCCTCCTT
12351	GCACTCAGCA	CCTCCCTTCT	CAATTCCTTT	TGCCATGTCT	CCACTTTAAA
12401	TTGGGTTGCT	CTCCAGATAG	TCTTTTCCTT	CACACTATTG	GCATCTGTGC
	AGAAGGTGCC	ATGADAGGAT	TCTCTGATCT	ATGATTTCTT	TGCATGATCA
12551	CGAACTAGCT	TCATGATAGC	ACCAGGAAGA	CTGATATCTC	CCACCAAACA
12601	AACCACTCAT	GGTGGTGCTC	TTTTTGCCTT	CACTATGAAG	TGTTTGTCTG
12651	CCTGTATGTG	AAAACGAGAG	GGTTTAATTG	TAAGGATGCA	GCACAGATTG
12701	GGACTGGCAT	CAGAAAGCCA	TTGGGGACTG	AGGTAGCTCT	AGAGACCGCT
12751	CATCTCTCC	AGTGCTCTCC	CTCCTGGGTG	ACATGTTTTC	TGTCTCCTGG
	TACCCCAAAG	TCTCTCTATG	GGCTTCTTTA	TTATTTGCAG	CTTGCAATGG
12901	CCTACCCACA	AAGCTCTTTC	TATTCTTCTA	CTGCATATAT	TCAACACAAC
12951	AAATCTGATT	TTTTTTTAAC	CTGGTCATGT	CAAAGACCAC	TGACCACATA
13001	TGAGCTGGTT	GCCCTGTGTC	AAGTGCCCCC	TTCTCCCACC	CTCTTCCCCT
13051	CCCCATCTGG	TCTGTCATAA	CTGAATGATG	GAGTGGGAAA	TTGAAATTGC
13101	CATGGGAATT	CCATGATAAG	CTATCTAAAC	AGTTTTATCT	ATAAGTGGTA
13201	GACAGAGTCA CCAAACTGGC	ACACATTCTA	AGTCCCAGGT	GAGACAGGCA	CCTGTCAACT
13251	TTTTGTAGTG	GCCTGTACTG	GGGCGGTAGG	CTGGAGAATG	AGAGCACTGA GCACAAATAC
13301	CCACTTCAGA	ATCCCCCAGC	CCAAATGCAT	CAAGCTCACT	ATAGACTCTG
13351	CAGCCACGAT	TCAGCTGGCT	TCTGCTCAGA	TCAACAGAAA	ACATTCTTAG
	TGAATGATGC	TTGTGGCACA	TATCTCAAGG	CTACCAGGGT	CATTTCTTCC
	CATTTACTTT TAATCTTCCG	TTCTCTGATC	TATCCTCTCC	AGGACACTAG	CGTCAGAAGA
			GTACACTATT ATTTCCAGAG	CACCOMORGO	GTCACTTTCA
			TTCATGCCTG	CCAGGGATAG	AATGACAGTA
	CTCCTGAGGC	TCTCCCTCCC	CACCCCTCCC	CTGCTGGACA	GCTGATCTGC
	TGGACTCAGC	CAGAGCCAGC	AGGCACCCC	TCTTTATCCT	AGGAGCTGCA
	AACTTGATGC	CTTTCCAGGA	AATCCCCAGA	AGCTGGAGTA	TCCTCATCTA
13801	CATGTGGCAC GACTGGGGTG	AGTGTATGGT	TGTGTCAGGT	GCTCATGTCC	CATTGCATAG
	AGTAGTGGCC		ACCGTCCTTT	AACCACTTCT	CAGTCAATG
13951	CTGCGCTCAG	TTGTAGATGT	GAGAAGAAAA	GGCCAAATAT	CTGCCAATCC
14001	TAGTCCTGGG	ATTCAAGATA	GAAAGAACTG	CATGGAGTGA	AGAAACTAGG
14051	AGTCTCCATT	TCACTGAGAT	GCATAAGAAT	GAAATTATTG	TCACTATTTC
14101		GGCCAATCCT	AATAAGAAAA	CCCTTTTTGA	GTCTCTCTTT
14201	TCTTTATCCT CACAGGGACA	ACATATAACA	CAGAAGCTTT	TTCTATTCCC	TGGATGAACC
14251	CAGACTAGAA	TCTTCCAGAA	GCACTGCTAA	CCTACTCACT	ATTTCTTTAT
14301	AGACAGGTGG	TTCTCAAGCC	AGCTCCCCAC	CGCAGGCCTT	TTTCATCCTC
14351	TGCCCCTCCC	TGTGGAACCC	ATGTTTTAGG	TTATTAGCTG	ATAATTGGAT
14401	TTCTATTTTT	TCTCATAAAA	TACAGCAAAA	GATAGCTAGT	GATATTATCA
14451	TGAGTTAATG	TAATTATAGC	CAAAGCAGAG	AGAAACAACA	TTTTAATTAA
14551	CCTGTGTGGA AGAGACAGAA	ATGAACACAG	CCAACCCCTC	TTCTATTTTG	GGGGTTGAGT
14601	TGATGTAAAA	TGCTTTGAAA	TTATTGGGCIG	CTCATTCTTT	AAACTTCTTTAAC
14651	TTGATGATGG	TAACTCCGTA	AGGGGATCAG	AACATGCTGG	AAAGAATGGG
14701	CACAGCTTTG	GTTACCTGGG	CCTTACCACT	GTTATTCAGG	CCTCTGAGAA
14751	AGCTTACTAT	TGTTGTTATG	TTTCTTACAT	AATAAAACTT	CTAATATTTG
14801	TATGAAAACA	TAGAATTCCA	CTTTTAAAGA	TGTAAGGATT	TTGTCATACC
14901	ATTAGGGTTA TTTACCCGCG	AACACAGAGT	TTGATTCTAG	GTCTAAGAAA	TATTAAGTAA
14951	CTAATACCAC	ATATTCTCAC	TCATATGTGG	GAGCTAAAA	TATTGATCTT
15001	AAAAAGGTAG	AGAGTAGAAT	TGTAGTTATT	AGAGGATGCG	AAGGAGGATA
15051	GGGAGAGGTT	GGTTAATGGA	TACAATGTGA	AGTTATGTAA	GAGGAGTAAG
15101	TTCTAGTGTT	TTGTAGCACT	GTAGGGTGAA	TATGGTTAAC	AGTAATTTAG
15151	TGTATATTTA	AAAAAAAAA	GACAGGATTC	TGAATATTCA	CAAAGAAATG
15251	ATAAATATTC GGGAGGCCGA	AGCTGGGCGT	GGTTGCTCAC	GCCTATATTC	CCAGCATTTT
15301	TGGACAACAT	GGTGAAACCC	CGTCTCTACC	ATABAGAGTTT	BABATCAGCC AAAATTACCT
15351	GGGCATGGTG	GCGCACACCT	GTAGTCCTAG	CTACTTAAGA	GGCTGAGGCA

FIGURE 3, page 4 of 57

			GAGGCAGAGG		
			GTGACAGAGT		
12201	AAAAAAAAAA	GAATGATAAA	TATTTAAGGT	GATAGATATG	CTAATTACCC
12221	TGATTTGATC	ATTACACTTT	GTATACATGT	GTCAAAATAT	CACTCTGTAT
			TATGTGTCAA		
			AAACATATGT		
12/01	TAAATTATAT	CTAAGGGTGT	GATAAAATTA	CAGTATAAGA	TTGTGCTTGA
15/51	AAAAGTGCAA	TAAGAAGTAA	ATATGTACAG	ATGAGAAAAA	GTGCAAAGAA
12001	CTAAGTCCTA	AGCAGACTAT	ACCTTTCCTA	CTGCATGGTA	CTTCTCTGGC
12821	CTTTTGCTTT	GAAAGATTTT	GCACCCAGCA	TGGCAAGTGG	TTAGCAGAGG
12901	CAGCCATTCT	CACTTGTGCG	TTGGCTTTGG	GAGCCATATA	TGTTGTTCAG
12921	CTGGGTGTGG	AGTGGAAAGG	CTGCATGTTG	TATTAATGCA	TTGTTAAGAA
			TGGGAAGTGA		
16021	GAAAGACAAC	TTTTAATCTT	TTACTTTACA	CTTTGTGCAC	TTTTAAATGT
			TTTAATAATA		
			AACTGCATAT		
			TATGAAGCTG		
16231	CCTCATATGT	AGGAAGTTAA	GAATGCATTC	TACGTTTCTT	CTTTAAGGAG
			ATAGGGGTAA		
			AATGCCCAGC		
			CCCACCCTCA		
			ATCACCAGCA		
			AGGGCGAGGC		
10001	AGATCAGGGG	ACCTCCTTTG	ATGCCATGTC	CATGGTGTCC	GAGGGCAGCC
			GCAGTGATGA		
			CTATGCCAGC		
			GCAGAGGGGT		
			GAGCTCATAC		
16051	BARAGGCACA	GTCCCTGTCC	TCAGGGAGGT	CACAGTTGAT	AGGGAAGACA
16001	AGCATATGTG	CTAGCTGCTA	TAGAAGGGG	AACCACTGAG	GGCTGTGGCC
16901	ACACAGAGGC	AACACCCCCT	TCTTGTTTTT	TTGTCAGGGA	TTCAGTTTGG
17001	CGICATIAGA	AGIGACTIGC	ACAACCCCCT	CCTCCAGTCA	ATTCAGAAGG
17061	CARCCAACRA	CAGGAATGAT	GAATTAGCTT	CAGCTTGTGG	GGCACACACA
			TCAGGAGTAA		
			ACCCGGTTCT		
17201	GCCTCACACA	CCACCAGGAC	ACAATTTATG	CCTGCAGAAT	GAATGAATGA
17261	MCCACCAACC	GAATTCCTGG	AACCTCTTCT	GCTTATGTGC	CACACCAGGT
			CTGGGACTGG		
17351	TCTCCCACCT	GCACCCACCA	TTTTGCATTC	TTCAGCCCTT	CCTCCTCTCC
17401	COCCACACC	TCAGCAATAT	CCACAGAGCC	CTCTGAGCAA	CTCTGAGCCT
			CTGGGCACCA		
17501	ACCCTCTAAT	TC A ATT A A CC	AGCCTGGGCT CTTCTCTCCA	GCCTTTGATG	CTCAGGAGAC
17551	TATCTCCCC	TURALITARGU	AGCCCCCTGC	GGGAGCATGT	AATTATGTCC
17651	CACATTCCAC	CCACAGAACCI	TTCCCTGGGA ACAGCACTTT	GGAAACTAAT	CTGCTTAGCC
17701	CHCATTGGAC	ACACAACCCC	CCCACTCTGG	CCCAACTCC	CCCCCCCCCCCCCCCCCCCCCCCCCCCCCCCCCCCCCCC
17751	CTCCTCACIG	CCTCTTTCCC	ATACAGATCT	GGGAACTGAG	GGCTCTCTTG
			GAGCTCCCCT TGGGGGTTAG		
17951	MAGACIGAAC	mcmmcmcccc	TGTTTTTGTT	CTTGTTGTTG	TTTGCCTGTT
	ACACTTCCCC	TCTIGIGGG	AGAGGGTATA TCATTTGAAT	MUNITUATIO	ACAGAGTGGC
18051	ATTTCTCTCT	TARCATCCTC	CAATTTCTCT	CCUTACCAM	MCMCMCCMAR
18101	CATCCTCATC	AATTCCTCCA	GGTGTTGGCG	CCATCTCCCA	NCTCTGGTAA
18151	СТСТТТСТСА	GTTTTGGGGGG	AAGTTGCCTT	DATIGIGGA	VALOTOCCIO
			TCTGTGCAAT		
18251	CTCAGGGAGG	TGTTGCTTCC	CACTGCCCAC	GCCACTGGAA	ALCACTACCC
18301	CAGGTTTACT	CGAGTCCTCC	TTTTGAGGAA	CCCAAATTCT	TOCAGIAGOC
18351	TTATGTGAGA	TCTGCCCAAA	ATGCCATTGG	CAAGCTGTAC	TECETTICIT
18401	AGTGTCCTTC	CTCCTCCCAA	ATGTATGTCT	DCTCCDDACC	ACAGGATACT
18451	ACCTTATTTG	GGAATAGGGC	TTTTGCAGGT	CTAACCATTA	ACAGGATACT
18501	TGAGGTTATA	CTAGATTAGA	ATGGGCCCTA	CATCCTATCA	CTCCTATCCT
18551	TACAAGAAGG	CCATGTGATG	ACAAAGACAA	DEDDTEEDET	CIGGIAICCI
18601	AGGAACTCCA	AGGATTGCTA	GGAAACACCA	GAAGCTTCCA	GGAACCCCA
18651	GAACAGATTC	TCCTCTCGGA	CCTCTAGAAG	GAATCACTCC	TCCTCATACC
18701	TTGATTTTGG	ACTTCTAGCC	TCCAGACCTG	TTGGGGAGAA	TACIGNIACC
18751	CTGTTTTAAC	CTACCACGTT	TGTGGCGATT	TGTCACACCA	THOUTITOIN
18801	ACTAATACAT	ACAACCTGCA	CAATGCCTAC	TCCACCAMOCA	CATACCAACA
18851	CAAGGGCCTC	ACABTTATOT	CCAAAGGACT	CATACARCAC	CATAGCAAGT
18901	GCTACTTCTC	CCTCAGGACC	CTGACCCACA	CCTCTCX XCC	CACCACTAT
18951	CCAGAGCTCA	TTCABCAACO	TTGTTATATA	GCCCTTCAAGG	CAUCHACTACC
19001	TTTCABTTC	TCTTTCCAACI	TAGATGAGGG	TTCARA A A A A	1 TGTAAACCT
19051	TTCTCTANCC	CACATACCCC	AATCTGTTTT	TIGHAAAATA	MATGGCCACT
19101	ATABTECCCT	CCTCTTCTDT	CTTCCAATCT	CCATACCTCAT	CCTTCCTTC
19151	TAGTTCTTTT	TOTOTIOINI	TCTTTTTTT	PCCCCC PCCC	TOTAL CTTGA
19201	GCCTGGGCCTG	GAGTGCAGTG	GCACGATCTC	CCCTCACTC	CACCECECEC
				OGC 1 CMC 10C	CACCICIOCC

FIGURE 3, page 5 of 57

	TCCCAGGTTC	AAGCAAGTCT	CCTGCCTCAG	CCACCTGAGT	AGCTGGGATT
19301	ACAGGCACCT	GCCACCATGC	CTGGCCAATT	TTTTGTACTT	TTAGCAGAGG
19351	TGGGGTTTCA				
19401	TCCACCCACC	TCAGCCTCTC			ATGAGCTACC
19451	GCGCCTGGCC	AGATAGTTCT	TAAACAACTG		
	CAGGGGCAGC				
		CATGAACTGC	ATTGCTCATT	TCTGCTTTTT	GACCTTTTCG
19551	ATGGCTGAAC	TCTAGGCCAT			
19601	GTCATTTTGT	GACTAGGGAG	ACAAAAAAGG	GCCTATTCTC	CAAATCCCCT
19651	TTCCCTCTGG	AGTTCCTCGG	TGCCTTAAAG	CTTGTCCTGA	GCTACAGGTG
19701	TGTTACCTGC	TTATCCCAAA	ATGCAGGCAT	GTTACCTGCT	TTCCTCTGCA
19751	AAGAGAGGCA	GGCCTGGCTG	GGGCACAGCT	GAAGATGTCA	AGGCCAACCT
19801	AAGGGCAGCC	AAGCTATGGC	TGTCTGTGAC	ANGAGGAGAG	CACCCCTCAT
19851		GAGGCATTGA	CTTCATCTCC	CCCMMMCCccm	
19901					CCTACCCTCC
		TGATGATCCT		TGATGAGTTC	AAGACAGAAG
19951	TITIGCCTCAT	CATTGCCACA	ATAAAATCAC	CAATAACAGA	AGTGTGAAAG
20001		AGTGGAAGCC	CATATATACA	CAGGGGGTAA	TAGAGCAGCA
20051	TGATTAAATA	TGTGGCCTTG	TTATCAGACA	GGCTGATTTG	GAGTCCCAGC
20101	TACTTGTTGG	TGACCTGAAC	TAGAGGAAGT	TATCTAACCT	TTCATTTTAC
20151	TCATTTACAT	AACATGGCTA	ATAATAGCAC	CTACCTTATA	GGGTTATTGT
20201	GAGGATTGAA	TACAATTATG	CAATATAAAA	CGTTTAGCAT	AGTGCCTAGT
20251	CTAAATTCCT	CACCAGGGGT	ATCATCTACT	ACTOTOTO CO	TAAGTAATTA
	GTATCCTCCA	CATGTCACAG	CCAMMMCACC		
	TCACCTTCCC	CATGICACAG	CCATTIGACC	TATCTGGGCC	AGCGTTTTGC
20351	TCAGGIICCC	CCAGCAGTAA	TTGTATTCCC	TCCCCAATCC	CGGGATTAGC
20401	TTTTAGGAAG	AAACAGTTGA	TCTAAAGATA	GAAAGTCAGA	GTACTGTCTG
	GAGGAAGGTA	GAGGGAAATG	TCATTATCTG	GGTTTTCTTT	GATGATGTCA
20501	GGGAACATGA	CAGGCTGCTC	CCAAAGACAG	AGCAGCCCCA	GGACAGGGAA
20551	GAAGGTGACC	TTGAGGTTGA	CTCCTCTGCA	TCCCGATGTG	GACGTTATGG
20601	ACTTGTTTTG	GAGATGAAGG	GAAAGAAAGA	TGGAATGTAG	AAAGTGAAGG
20651	AGAATAAAG	AAGTGGGAGG	AAGAAGGGCT	GGGAGGAGGA	TGGGCAAAGT
20701	CTTTCTGGTC				
20751					CAGTGGGACT
		AGCAGCTACA			AGAGGGCTCT
20801	TGGGCATGTC	TTGGAGCAGC			TCTCTCACTA
20851		CCAGATCACA		ACCTTCCACC	CCCGGGCCTG
20901	TTAATGATCA	AAAAGCTTAA			GAGTGGAACC
20951	ATATCTCTGG	GCTCCTGTGA	TGAAAACCAC	AAGCCTGTCA	GGCTGGGGCT
21001	GCTTCACATG	GAGGGCCCTG	CTCTTAATGG	CCAAGTGATC	TGGAGCAAGA
21051	CCCGTGACTC	TCCCATAGTG			CCCCACGCAT
21101		GGAAGTTCAG	TAACTAAGGA	ስጥጥልስርጥስጥጥ	CTCCAGCCTG
21151	ATTCTGCTTT				
21201		AGTCACCCTT			CATCCCTATA
21251					
		ATGAAGTGCC		GTGATTTATT	TAGTACTTAC
21301	TGTGTGCCAG				GTATGATCCT
21351	TACACTAAGC			CCCTGACCAC	TCTGTGCTTC
21401	CCTTTTCACA	ACACAGCTTG	TCACTAAATC	CAAGTCAGGA	ATTCCAGGTT
21451	AGGCTTGAGT	TGTGCAGAGC	CCTTAACTGA	AATTTGCCAT	GGTTGAGGCA
21501	TGATTGCAAT			TCTACACACC	TACTTGTCAT
21551	ATTCACGCCC	TGATCACGGC			CTTTAGAACT
21601	TCTTTCCTAT	AGAACACGTT			
21651	CCTGGCCTAA				ACTGATCAGC
21701				TTGCACATAG	CTGGTTGAAT
	CGTATGTATT	GCTGTTTGTG		·-	AATATCGGCA
21751	ATTTTATGTG	TTTCATTCAA		CCAGCATTCT	TACCTTGTCG
21801	CTTTGTAAAC	CCTGCTGCTC	TCAAATCTCC	ACTAGCTGTT	TCCTGAGCAG
21851	AAGGAGATAA	AAGGCTGGCT	CACACCCCCA	TGTTTTTACT	GGTCACAGTT
21901	ACTGCCACCA	TCCAAGGCTG	AAGAGACTTC	CTTTGTGTTA	GGGCTAAAAC
21951	CTTAGTCATT	GTATCTAAAT	GTCTTCTGTA	TTCCTTTCCT	CAAAAGAAAA
22001	AAGTACCCTC	TTCTGCCAAC	CCTCTCCCAT	GCCAACTAAA	CAAGCAAGCA
22051	AGCAAACAAC	AAAGAAAAGG	TGATATTACA	GATGCTGCTC	AGCCTATGAT
22101	GGGGTTACAT	CCTGATAAAC	CCATCACAAG	GGATGTAATT	CCATTCCAAC
22151	TTACADATAC	CATAAGTCAA	מאחרתי מיינים	Thurbury war	DOCCACACA :
22201	CCTCATACCT	TAGCTTAGCC	MANIGIATII	MITTCATATA	ACCCACAGAA
22201	COLONIAGE	CCRCRRRR	TACTIGATCA	TGTTCAGAAG	ACTTATATTC
22221	GTCTACAAGT	GGACAAAAAC	ATATAAAACA	AAGCCTATTT	TAAAATAAGG
22301	TGTTGAATAT	CTCATATAAT	TTATTGAATA	TTGTACTGAA	AGTGAAAAAT
22351	AGAATGGTTT	TCTGGATACT	CAAAGTATAG	TTTCTACTGA	ATGCATATCA
22401	CTTTTGCACC	ATCATAAACT	TCAAAAATTG	TCGGTCGAAC	CTTCCTGAGT
22451	CAGGAATCCT	GTCTGTACAG	GGTATAAAGG	AGGAAAGCAT	CAGCTTTGGA
22501	GGCAGGTGGA	CCTGTGTTTG	AACCCTGATT	CTGCTAGAGC	TTĠACAATGC
22551	ATATTCGTTT	TCTATTGCAT	AACTAATTAC	TACADACAAC	<b>ፓር</b> አጥጥጥ አጥጥጥ
22601	СТСАСТТТТС	ATGAATCATG	AGTCCACCCA	CDAMMANA	CCACHMARACC
22651	TETTACCTCC	GGCTGCTGTC	TO TO CHOO PACE	CAMCCCCCC-	GCAGTTAAGG
22201	TOITMOUTED	ACCACCACACA	TIMICIGAAG	CAIGGGGTG	GGGGTGTGGA
22751	TTCCAAGGTC	AGGTGGTTGT	IGGCAAAATT	AATTTTCTTG	CAGCTATAGA
22/51	ACTCATGGCT	TGCTTCTTCA	AGGACACGGG	GAGAGAGAAT	CTCTCACATC
22801	TTTTAAAGGG	TTCACCTGAT	TAGGTCAGGT	CCACTCAGGA	CAGTTTCCCT
22851	TAAAGTCAAG	GCTTAATAGT	CAACTGATTA	GGGACCCTAA	TTATATCTGC
22901	AAAATACCTT	CACCATTGCC	ATGTAACATA	ATCATGGCAA	ATAATCACAG
22951	GTCCCAAATG	TTCACAGGTC	CCACTCACAC	TTGAGGGAGG	GGATTATATA
23001	GGGCATGTTC	TTGCGGAGAG	AAGGAATCTT	ACAGCCACAT	TGGAATCTCT
23051	CTTCCATGCT	ATTTGACCTC	AGGCAAATTC	ACTA ATCTCT	TCAACCOMC
					LONAGGITCA
		~	TY CY TO	~ ~	_

FIGURE 3, page 6 of 57

23101	ATTTCCTTAC	CTGGAATAAA	AGGACAATAA	GATCAGCCAT	ATAAGGCTAT
		AAATGAGATA			
23201	GACACAAGTA	TATAACAATT	TCCCTCCTAC	TGTTCCTTTT	GTTTTTCACC
23251	TATCCTGCAG	TCTCTGTCAC	TTCAAATACC	ATAGAAAACC	TTTCCAAGCA
		TGCCCCCAAA			
23351	GTTCCATAAA	<b>GTTAGCACAA</b>	ACTCCGAATG	AGTGAATCCT	AAAGCGTTGC
23401	TCCTGGAGGA	AATACAGGCT	GCTGGTCACA	ATATTTTTAT	CAACTGATCA
		TGTCTTATGT			
		ATTCATTAAC			
		ACAAAGCTTA			
23601	TCCCAGCCTT	CTTGCACTTA	GGATTCAGCA	GTATGCTTAA	GGGCCATTTT
		CTCATCAGCG			
		CAAAAAGGAC			
		ATCACCTTGT			
		TTCAAATTTT			
		TATTTTTTTC			
		AAGAGCTGAA			
		CTGGGCTTCG			
		CTAGGCATTG			
		ACTTCATACT			
		ACTCAGCCCC			
		ATTTGGCTGG			
		ACTGTAATGT			
24231	MCMCMMMMMCC	TTCAAGTGGT	TAGAGCTGGC	TACGGGTGGG	CTGAACAAGA
		TTCATTTCCC			
		CACTTTGAAG			
		ATCATTTGGA			
		CCTCCCTCCC			
		ATCTGCTTTA			
		AGTGCCATTG			
		AGTCCACAAC			
		AAGGTAGGGC			
		GTTTATACGT			
		CCCAAAGGAA			
		GGGCATCAAA			
		CAAATGGAGC			
		TCTAGTGAGG			
		CAGACCTGAT			
		GCAGGCACCA			
		TGTATTGAAT			
		GAAGGTTTAG			
		GGTGAGGGAT			
		AGCATTTACA			
		AATTTTCATG			
		TCATTTCATA			
		AGGGCACTAG			
		AGCTGAGCTC			
		TGAATGTAAG			
		CATCAAACAA			
		GAAATTCATG			
					AACGGAAGAA '
25651	TGCAACTGGC	AACTGAATGA	TATAGGTTGT	GATGACTGTT	AAATATCATG
25701	AAAAGAGACC	ATGATGAGCT	GAGGCACTCC	AAGAGACTTC	TTTTTGGAGA
25751	TATGTTTGGA	GCCAAATCTT	GAAGATTTAA	TTGCTTTTTT	CTTTTTTTT
		GAGTCTCGCT			
		CATTGCAACC			
		TGAGTAGCTG			
		ATTTCTAGTA			
26001	GGTCTCAAAC	TCCTGACCTC	AAGTGATCTA	CTCGCCTTGG	CCTTCCAAAG
		ACAGGCATGA			
26101	TTAAAAAAAA	AAAAAAAAA	AACAGGAAGT	TTTCGTTAGT	TTTTTTGTTT
26151	GTTTTACTTC	CCATAAAAAC	TCTTTGTGTC	ACATGGAGGT	GAATGGAAAG
		GCAACAGACG			
		AATGTATGCT			
		GCCCCACAG			
26351	CAGTCCTCAC	GTAGCTCACA	ATCCAGTGGA	GGAGACGGAC	TCAGAAACAG
26401	ATAGAGATGA	AGCCATGAGA	TCAGTACTGT	CCGAGGCCAT	GGCCACGGTT
26451	TTGTGGGAAC	CCACGAGAGG	GAATGACTAA	CTGTGGGGÄÄ	GAAGAGGGAG
26501	AGGACCAAAA	TGCAGGGGAA	GTGCTCACAG	AGGATAAGTA	AGCAGTGAGG
26551	TGCCATGAAA	TGAGTATACA	CCTGACAGCC	GTGTAACAGC	TCAGAGCCTG
26601	GGTAGAGGGG	AATAGAGCTG	CTGGTTCTCT	GGGGGGAAGA	GAGGGGTATG
26651	GGATTCTGGA	ACAGAAGCAC	CAAAACCAGC	AGGTTATTGG	AGCTGTTAGT
26701	GCTCAGATCA	GCAATGGGTG	CACAACCAAA	CCATTCTCCT	AGGGATGAGT
26751	TCTTTCCTGT	GGATGAGGGC	TTCTCAGCCT	GGCTTCTCCC	GAGAATTACC
26801	CGGGAAGCTT	GAAAAGTACT	GATGCCTGGA	ACCTACCTCC	AGAGAGTTGG
26851	ATTTCATTGT	GTTGACGTGG	GGCTGGGATA	TCAGTATATT	GTTTAAGCAC
26901	TCCAGGTGAT	TCTGATACGT	AGCTGTGATT	GAGAACCCTT	GCCCTAAGCT

FIGURE 3, page 7 of 57

26951	ATCCATCTGC	ACTCCAGGGG	TGCTCCCAGG	CCCATCTGTT	TGTAAATGGA
27001	CAGGTGTCTT	GAGGTAACAA	ATGTGCCAAG	GCTCTGGAGC	CAAGCACGCC
27051	TGGCTCCTTA	GTGCCTACTT	AGTGACCTCA	GGCAAGTTAC	TANATCCCCTT
	AAACTTTACA	AATCCTTAAT	TTCTAAAATC	TECECONATES	THANTIGOUIT
27151	TCACACCATT	ATTACGAGGT	TIGIAAAAIG	TGGGCAATGA	TAGTACCTCC
	CACTTCCACA	CCCCTCATCC	COMPCOCCO	TACTOTOAGO	TCATAATAAG
	CACTIGUACA	GGCCTCATGG	GCTAGGCCCT	CAAAACTTAA	CGCATCTACA
27251	GGCAACAGCC	ATATGAAAGG	AATTTTATAC	CACCAAGTCA	AAAAATCTGT
27301	GAGCACTGCT	CAGAAGCAAA	AGCCTGTCTC	CAACAGCGCT	CATTTAAGGG
27351	GTGGGCGAGC	TACAGAGAGA	AGAATGAGCC	CCCACAGGGT	AAGCTGGGGA
27401	AAGCTGGGGA	CAGAATGAGA	CTCAGGAAAT	CACTTGAATA	TTGATTATAT
27451	TTGTGCTCAA	TAATAAAATA	ACGAAATGAG	TACAGCCCTA	GACCTAAACA
27501	TTGTGGGTGA	GGCAAAGGCA	ATGCGTTAAT	TTTGCATCCA	CTGAGGAAAA
27551	ACTCTAAAAC	GGTGACTTCT	TTTTTAAGGG	ACCAGAAGAA	TCTACATTAT
27601	ATTTAGTCTA	AGTCAATACA	TACGACAGAA	CCTTCCCCTC	TAGACTTCAT
27651	AAGAAAGAAG	TAAAATAAGA	GAAAGAATAA	AAAACCCTTC	CACCAAAATA
27701	CTAACATTCA	GATAATGACT	TTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTT	CTCTCCCTTC	CACCAMAMIA
27751	CCTCAGAAAT	GAATAGATTT	CTCTTCTTAG	CCAAMCAMCA	GAGGAGGTTC
27901	CATCCACTTA	AGTGTGATCC	CICITCIAGI	GCAATCATCA	AAAGGTAATG
	TCCCTTTC A	AGIGIGATCC	CCAAGAGAAA	ATCAATGACC	TTTCTGTGTT
27851	TGCCTTTGAG	AAAATCAGCC	AGTCTATGGT	TAAATTAGAC	ATATTTTTC
27901	TCCTTGGTCA	AGATTAGTGG	GACCAAGAAT	GCAGTCTTAC	ACTCCTTCTA
2/951	GCAAAGAATT	ACCTGATGCC	TTATTTCACA	CAAATTTGCA	<b>AAGTTGTATG</b>
28001	GACGTTGTAT	CTTATTTTAA	GGAGAACTGG	TGATCAAATG	<b>ATGACTATTT</b>
28051	CAATAGTGGT	TCATTTACAC	CACCACCCTC	ACCCCACATC	CTGCTTTCAC
28101	CTGAATCTGA	ACGATCATAG	TCAGTCTGAG	ATTCTGAAGG	TTTGAAATTC
28151	CTTTTCTGAG	CTCTGCAAGA	ACAGCATCTC	CCAAGAGAGC	TCAGGGCAGA
28201	CTGTCTGGGA	GAGATTGGAA	ACCTGTCTTT	TGCAGTAACA	TGAATTGGTT
28251	GAATGGTCAC	CCTCCATATC	AGGCCTGCTT	CTCCCATTCC	CTTTCTCATC
28301	AGCCCAACTT	GGGTCTCACC	CTTCTCATTT	CTCCCATIGG	CCTCACATC
	GGCTGCACTG	GCCATTAGGT	CCCACCCTTC	CICICICIG	BCTCACATGG
28401	CACCTCCCCT	CTGTGGAGCC	CONNECTIO	COMOMOGRA	ACCCATTGGC
20401	CCACCCCAMM	CIGIGGAGCC	CTAAGGCAGG	GCTCTGGTCA	CTGGTGAGAG
20431	GGAGGCCAII	GGAGTCACTG	GGGTGGACCT	ACAGACCCTA	GGGTTAACAG
28201	CTAGGTGGGT	GTCCTCTTCA	GAGAAACGGG	TTACAAAGTG	AAAGAAAGTT
28551	ACACTGTGAG	GTCAGCCAGG	GAGGAAGACA	GAGAGCTGAT	ATAAGATAGG
	TACTGATTCC	CTGGGGATGT	GAAAGGAGGG	TAATATTCCT	AAAATGATAG
28651	CATTTAGCTT	CCAGTATACA	TTAATTGATT	CCTGATATTC	ATTAAAACTA
28701	AACGCTATTT	CCTTGATGTC	TCATCCAAAG	CCGCACCACT	CTTCCCACTA
28751	AGTCTGAGGG	GAGCTTGTTT	TGTTGACAAG	TGTAAGAGGT	TGAAGAGGGA
28801	CCCATGAACT	CTTTTGTCCT	ACTGAAGAGA	TCCACAGATG	GARACARATG
28851	CTCCTACCAC	ATTTATGAAC	TECTECTTE	CAGTCCCGCT	TCTGCTATCA
28901	TGCACAGGAA	CTGACTAAGC	TCCAAAGCCA	CACCATCTAA	ATCTCCCTCT
	AATAAATGTA	AGTCATTTAT	TACCTACATA	CACTTCACCA	ACTCACCTAL
29001	CCTGCAAATT	TCAAGCATGT	CANTCUTCON	TOTTTO	CCMACCIAA
	ACACTTTCAC	AAATGTATTT	A A TO CO COOM	TCTTTCATGT	
20101	CARECCERA	AAAIGIATIT	AATGTCTCTT	TGCTTCCTTT	TCTACCCACA
20151	CAATGGGTAT	AATAATGTCT	ACCATATATC	TTTGCAGCAA	GGTCTAAATG
29151	GGGTGATACA	TGCTGAATAC	ATTTCCAACA	GAGTCTGTGC	AATGATAAGC
29201	TCTTTCCAAA	TGTTAGTTAA	AGCTAACCAA	CTAACCCACC	AACAAACCAA
29251	CCTCTTAGCC	AGGACTGATG	GAAGGAGTCT	GTGAGAGAAT	GCATTTAAAA
29301	CACTTGGCAC	CATGCCTGAC	AAGAGTAAGT	ACTCGATAAA	TCAGTTATTG
29351	TTATTATCGC	ATCGGTATTA	TGACCATTAT	CCTCTTCTCT	ATAGGCTTCA
29401	GGTTTTCCTG	TCTTTTTATC		TCCAGCAGAA	
29451	TAACTAAGTC	TCTACTGTGT			
29501	AAGTGAGAAT	TTGGTCCTGC		CTTATAGTCT	
29551		ATAGAAAGGA	AACTAACAAT	ATGCAAAAGG	ANACTONTAC
29601	ТТТСТССТАА	ATGCCAGGTG	CTCCTCATAC	TECETTERES	CACAMCMCAM
29651	AGATGCTATA	GGAGGTCAAA	CONCANCECT	CCACCTTCAGA	CMARCHER
29701	ACCCADANCC	GTGAAAGAAT	TACTICATION A	MCMACA COMA	CIAAGITITC
20751	CACACTCCAT	CLGWWWGWWI	TAGICATIAA	TGTACACCTA	CATTACCTGC
20001	CAGACICCAI	TCAAAAATAT	TCTTACCAAA	TCATCACAAT	ACCTTGTTGG
23001	TAGGTACTAT	TACTATTTTA	CAGAGGAGGA	AAGTGAGGCA	AAGACACATT
29851	AAATAATTTT	CCCAGAATCC	CAAGGTGTGA	GGTGGAGCAA	GGACACAAAT
29901	CCATGGCTCT	AAGTCCCTCC	TAGTATATCC	TGCAAACACA	TCTGGAATTA
29951	ATGCAGAGAG	GAAGGGGAGA	GGCAGTGTTC	TGCAGGAGTT	CAGAGCCATG
30001	ATAACCCTTC	TTGTGTGGCT	TTTGGTAAGT	TATTTTACCT	CTTACCCTCT
30051	GTTTCCCCAT	CTGTTCAATG	AAGGTTGTAT	ATACACACAT	TATATGGCCG
30101	CTGTAAGTGT	GCAGTGATAT	GATGCATGGG	GACTCAGTTC	ATGAGGCAGT
30151	GTGAATTCTG	AAGGTATCAC	AATGGGACAG	GTGTTTTTT	CTCCACTCAT
30201	TTTCTCCGAA	AGTCTTTTGT	TTTGTTGCCC	TCCCTCTTTC	GGGCATATCC
30251	TTTCACCTCA	TACCTTAATG	DCDTCDCDAM	CTCCTCTTTG	CUCCCAACEC
30301	TTGTGGTTDA	AATTATTCTG	CCCMMCCAMM	UTGCMATITE	LIGGCAACTT
30361	CADADACCAC	CANADACS CO.	ADARA SER	TRAAGCACT	AATAGCAAAG
30404	GIWIINGGIG	CAAAATGATG	ATAAAAATAA	TTGCAATTTT	TACCATTAAA
20401	MGTCATGGCA	AAACCACAAT	TACTTTGGCA	CCAGCTGAAT	ATTTTGAAAC
20521	COCCTACTCT	GATGTTAACC	AAGTTCATGA	TTCAAAGAAC	TTGCAGAGGG
30501	GTAGGGGAAT	TTCAAGGGAA	AGGGGGAGAT	GCCTGGGGTT	GTCACACACT
30551	CTGTCTTTCA	TCCTCTATTG	ACATGTTGGT	TATTTGGAGA	TGGTATTCAG
30601	TTCCACTATA	GCCCCTCAGT	CACTGTAGAC	CCTCTCAAAG	GGGCAATCAT
30651	GTTTCCCTTA	GGTCAGGTCC	ATTCATCTAA	CCCCTCTCCC	GGGGGCATCA
30701	CCTTGTTTGT	TCCAGCAGCT	GTCTGGCCAA	ACTCACACCT	CCTCCTCACC
30751	CTCTAGCCCT	TATGATCTGC	TTTGGGGAGC	CATGGGAACC	CCTAGTTTCC

FIGURE 3, page 8 of 57

	TCTTTCATAC				
	CATTGCCTTT				
30901	GCTCCACACT	CCAGCAGACC	TTCTGCTGGG	CGAGAAGCTG	CAGGCCTGAA
30951	TCTCTGTGTT	CTCATATGGC	CCCAACTCTT	GGGATTACAC	TAGCTCTTGT
31001	AAGAACTCAA	TGCTCTGCTC	TGCTCATTTT	GATGCCATCA	AAGAGGGCTT
	GCAAGTTACC				TAGAGGTACC
	CCTAATCTTT				
		GGGCACTTCT			
31201	GCCAAAAATC		TCAACAACTC		CCTTCTCTCC
	TCTATTTTAT				
		GCCTTTTGTG			ATCCTCAACT
31351	GCTGGCTTCT	GTCCTTAAGC	CTGGGGAGAA	TTAAGTCCTC	TTTGCCTCAG
31401	TTTGGCACTC	CAATTGCCAA	CATTGGGACA	GCAGGAAAAG	TTCCATCCAA
31451	CATCCCATTA	AATATGTAAT	GTGTATTAGC	ACAGCGCCTG	GCACTGGGCA
31501	GGTATTTTCT	AAGTGATAGC	CAATGCGAAG	CCTACTTTAT	TATTTTCCTC
31551	TTTGCTTAAC	CTACAAGGTG	TCTAAGACCA	TTTGTTTGTC	CACACATAGT
31601	AAGATAAACA	GCACTGAGAC	TGTGGTCCTT	TCTGCCCTGT	GTCCTTATCC
31651	CACCTGGGAA	TCTGGAAAGC	CAAGCCTAGA	CACACTCGTT	CCACAAATGT
31701	TTACTGAAGC	TTGTTCTATT	CAAAGCACTG	TACAGCTACA	AAGACCATCT
31751	TTTCTGAACT	CCAAACCAGG	CCACATGGTT	GGAATAACTT	CAAGTATGGA
	GACCAAGAGA				
	CCAGGAACTT				
	AATCTGCTTG				ATGAGTTCAG
	AGCAGGAAGA				
	TTTATAGTCT				TCTTTCTTTC
	CTTTGGTCAA				
	CTAGGTACTG				
	GCGGAGCTTA				TGGCTGCTCT
	CTTTATTTGA				
32251	CTCCACAGAG	G.A.A.A.T.	TATUALISATE	CTCTGCCAGC	CACCCACCCA
	CTCAAGACCA				
		TTTCTICTUA			TCTGCCATCC
	ATTTTCCCTC				GAGATTGAGC
	GGAGTGATGG				TGACATGGGA
32501	ATGAGGAGAC	TTGCTTAAAG	GATAAG/TCAT	GCTAAGTCAT	CCATCGTTCT
32551	CCCCTAAGGA	GGTGAATTJA	ASTICCEATT	TTTCCCAGGG	AGCCAAATTA
32601	ACAAGGTGCT	GGGAGATTTC	CAAATTAGAA	AAAAAAAAA	AAAAAGGCAC
32651	CACCAGCTCT	CAAATTAGAG	ASSETSTTGA	GTTGTTTTTT	GGAGCAGATC
32701	ATTGTATTTG	GCATCTAACC	TTSAAATAGA	GGAGAAAGCA	TGGAATTTCT
32751	GCTGAAAACT	CATCCTTCTC	TGA /CAGGTG	GTACAAATAA	GCATCGTTGT
	GTTCTCAGAG				
	AACCACAGTG				
	TTTTACAATT				CAGCTACCCC
	AGTACTTTTG		TGCTTATAAA		GCTTGGCACG
	GTGGCTCACA				GTGGGGTGGA
	TCACGAGGTC				GTCCCTACTA
	AAAATACAAA				
	GCTCGGGAGG				
33201					
	AGAGCAAGAC	GTGAGCCAAG			CCTGGGTGAC
				AAGATATATA	
33301	TATAATATTC		CTAGTAGAAA		GTTTTTAGGT
33351		TGGTTCCCAG			
	TTACCTACAT				
	TGCTAGCTTT				
	AAGTAGCATG				
	GTGAAAATGC				
	GTGAGCCTGG				
	CTTGTATTCT				
	CAGTTTTAGG				
	AACCTATGCA				
	TTAAAGCAAA				
33851	GAAGAACAGT	GTTTTATGGG	GAAGGACTAG	TACACACAAA	GGCTGCAAAG
	GCGAGTGGGC				
	GAGAGGGAGT				
	TCACGCAGGC				
	GACTACCGTA				
	GTATTGCCTC				
34151	GGGCTGCGCT	CCCTCCAAAA	CCTGTAGAGG	AGAATCCTTC	CTTGCCTGTC
34201	CCTAGCTTCC	AGTGGGTTGC	TAGCAATCCT	GGGCTGCGTC	ACTCCACCTC
34251	TGCCTTGGTT	GTCACAGGGC	GTTGTCTTTC	TCTCTGGGIG	DCTTCDCAGCIC
34301	GCCCTCTTCT	<b>ጥርጥጥርጥጥጥጥ</b>	GTGTGTGTCT	CTCTCTCTCTC	DCTCTCTCTCT
34351	ACAGAAGTTT	ተተጀመተያቸው ተ	ATTTATTCE	THE THE TAIL THE	CAUTCAUACA
34401	CATAATAGTT	PACCEATIVITIE	TTCCCCTACA	TIMITITATIT	CATIGATAAA
34461	TACACTCTCT	CATARTAGTT	TCACCCCCC	TGAGATATTG	DATACATGTG
345V1	TACAGTGTGT	TC A TOTAL CARA	CAMPAGGGGGAT	TGGAATATCC	ATTCACCTCC
34501	AAACATTTTC	ACARTTUTT	ATTGGGGAC	ATTATAATTC	TTCTAGCTAT
34501	TTTGAAATAT	CARCANTAGATT	ATTGTTTACT	ATAATTTCCC	TGCTGTACTA
7400T	TCGAATACTA	GMACTTATTC	CITCTGTTGA	GGGTGTACTT	TTGCACCCAT

FIGURE 3, page 9 of 57

34651	TAACCAACTT	TTCTTTATGT	CCTCCTTCCC	ACTTCCCTTA	CCAGCCTCTC
34701	GTAACCACCA	ATCTACTCTC	TACCACCATG	AAATCAACTT	TTTTTTTTT
34751	TAGCTCTCAT		GACTATGCAG		
34801 34851	CTTATTTCAC TGACAGGATC	TTATTTATTT	GACCTCCAGT		
34901	TGTATATCAT	ATCTTCTTTA			
34951	GATTCCATAT			CTCCAATAAC	TATTTAGGTT CATGGAAGTG
35001	AAAATATCTC	TTCAACATAC			ATATACCCAG
35051	TGGTAGGATT	GCTAGATCAT			TTTTAAAGGA
35101	ACCTCCATAC	TTTTTTTCCA			TTCCCACCAA
35151 35201	CAGCATATGG	TCATCTCCTT	_	CTTGCCAGAA	TTTGTTATAT
35251	TTTGTCTTTT TGTAGTTTTG	TGATAATAGC	CATTCTGACT CCCTTATAAT	GGGGTAAGAT	GATATATCAC
35301	TTATATACCT	GTTGGCCATT			GAGCATTTTT
35351	AGGTCTTCTG		AGTGGATTAT		GCTACTGAGT
35401	TCTTCGAGTT	TCTTATATAT			TTATGAGGAC
35451	TCCAGTTATA		GAGGTCCACC	CTTTTTCAGA	
35501		ATTACATCTG		ATTTCCAAGT	AAGGTCACAT
35551 35601	TCTGAGGTAC AGCTCAACAC	AAGGGTTTAG		TATGAATTCC	AGTGGGACAC
35651	TGAGTGTCTT		GGTAGGGAAC ATGGACTGGT	TTTATTCTAC GTGATGTATG	TTGCAAGTTC
35701	CGCTGTGTGA			GGATGGAAGT	TGGAGACTAA
35751	TAAAGGACTA			AGGTGAGAGG	TGATGATGGC
35801	AGAACTAAGG	TGATAGCAGT	AGAGAGAAGA		
35851	ATCTTTTGCA			CTATGGATTG	GACATGGGAT
35901 35951	GAGGAAAAGG			TAGGCTTTTT	ACTTTAATCG
36001	TGAAGGGAAG		GTATGGACGG	CGGACAAACC	TGGAGAGGAT
36051	AGGGATTAAA	AATTGGAAAT		AGTCAACAAT	GGAAGAATGC ATCTTACTTT
36101			AAAAAGCATC	CTTTTGTTGG	AAAGCTCAAT
36151	CCTTGTTAAA	ATGAAGACAT	CTCTGGGAGA	GGAAACATAG	TGAGCACCTT
36201	TCCCAAAAGC			GACAGAGTAG	CATACAGGAC
36251 36301	GTCTTGGCAT		GACAGAAAGA	GCAATGTAGG	ACAAGGCAGT
36351	TTAATTCCAT	CACAGTCTTT	CCTCCGACTG	GCTGTGAGCA AGTGCCCAAA	AGTGCTCAAT
36401	CAAAAGTACC	AGCATGATGG		TAGCAAGTTC	CCTCCACAGA
36451			AGAGGGGAAG	TTGACCCCTG	GGGATGGGGA
36501		AGGAGAACAT	GAAACTGAGA	AAAGGGCTTT	GAGTGAAATC
36551	TAGGCTAAAA	GCTAAGGTTT		CCACCATTGA	CCCAACATGA
36601 36651	CCAGGGCTTT ATTCCTGGAA			GATACCCCAT	CTTCTTCTGT
36701	TGGGTTTTCA	CTAGCTCTCC		AATTGTGCTT TGTATCTCTG	CTATCAGAGC
36751	TTGTTTGAAT	TCCTGCCAGG	TCAGCTGAAT	TTGGGCATTT	TTGCCCTATT GGGGTGAAAA
36801	ACCATCAAGT		GGCTTTGGCA		TGTGACCCCA
36851	CTGGTCTCTC	CCTCACATTT	GCTGTGGTCC	GTGCACGGAA	TTTGTCAAAA
36901	GACCTCCTCA	GTATCAGCTT		TCAATGCACC	TTGTTCTGAA
36951 37001	TAGGATATTA GGGTCCTTAG				ATGCCAGAAG
37051	TGGCTGCAGA		GTTTTTTCTG TGTGGACTGA		AAGGAGGAGG
37101	AGGACCATTT				GCCTGGTGTC CTAAACTGTA
37151	AGCACAAGAG	AGAGTTCAGC	ATCATTTGCA	TCCTATTTTA	TTGTCTTTCT
37201	TCTCTTTTCT	TTCAAGGCCT	CATTTTTTT	GGCTTGAACA	AATGGTAAAG
37251	GCCATTTTAT	TACAGGTACC	AAGCCAAACT	TTCCTTGGTT	TTGTGGCCAT
3/301 37351	TETTTEGTET	CACCCCCTCC	TCCTTTACTT	TAAATAACTT ATTTTCTAAC	TAAAAACATC
37401	TTTATGGGGG	TGTTTTTGGG	GGGGTTTATT	GAGTGTCAAA	CCTCCCACTA
37451	AATTAGAATC	AGAAGACAAC	AGTTAGTGAT	AAGCAGAGAA	GCCAAGGATG
37501	TTACCATAGG	CAGGCAGCAG	AGAGAGGGGA	ATTGGTGGCT	GGCCCCCAA
37551	AAACAGATTT	GAAGATCTCC	TTCTGTCATG	TAGTGAATCC	CCAAGTGCCT
37661	AGGGTGGGCT	GTGATTACTT	GAGCTCCTGT	CTCCACTGTC	
37701	TGCCTTGGGG	ACCTCA ATTE	ACACACATTT	GCTCATAGCA TTTAGATACC	TCAGGTATTC
37751	CTTTTAACAC	CAGATTGCCA	GGATCATGAC		TACCCTGAAA
37801	TGCAATTGAC	AAATGGGATG	AAAGATTTCC	CGTTTCATCC	ACATTTGCCT
37851	CCTGAGCTAC	TTACAGCAGC	AGGTCACCGC	AGCCAGAGCC	CACCTGCTTG
37901	CCCACCATGC	CCGCACACAG	ACAATGCTGC	TTCTGTGGCT	GGAGGTCGGA
37951	ACACCTCAGC	ACTATCTCAG	TTTGGCTGCA	GATCCTCTGT	GTGCTTGGTA
38021 2000T	AACAGGTTTC	CTCATCTGTA	AAATGAATTG	GCTCTTCCAC	AACTTTTTTA
38101	AATACTAAAA	GAGTGCAAAC	GGATGGGGTAA	ATACTCAAAT CCAAATATTA	GCTAAACTCA
38151	TGCAGCATTT	TCTGACCTTG	CTGCTTTTTC	TGGTGAGTGG	CAGTGAAGGC
38201	TTAGTTTGGT	TTCTTCTCTC	CCATTCTAAT	CAAGCAAGAA	GTGACCACCA
38251	AAAGGGGCAC	TCACCAAACC	AGAACAAGCT	AGTTCTTTCA	TCTTTAATTC
38301	ATTGCAACCA	AACAGATGCC	ACAGAAAGAG	CCAAGGGCTC	CAGGCTTTAG
38351	CTCCAGCCTT	GCCATTAACT	ACATATGTAA	GTCAGCCATG	CTGGTCTGCA
38461	CTATTCCCCC	TIGCATGATC	AAGGGACAAC	TTGGAAGGTC	TCCAATCACT
~047I	CIMITOCOCO			ATTTCCTGGA	GATGTCTGTC
		7	TOTTO		4.0

FIGURE 3, page 10 of 57

38501	CTCCTCCCAG	TTAAAGACAG	ACCTTGACCC	ACCTCCACTT	CCTTCTCTGT
38551	GGCCCTGTCT		CTTGTTCTTG		TGTTCTCTCA
	CCGTGTTTGT		CTATCACTGT		
					ATTGTTTTTC
38651	TAATGTCCCT		CTGATTTTCA		
	AAACACTTAT		ACATTCTGTA		TTCCCTTTAT
38751	CAAATGCAAT	CTAAGAAGCT	CACAGTTTCT	CTCAGTTTCA	ACAAGAGAAA
38801	TCAGGAGCAC	TTGAATTATA	CAACTTGACA	TTATTAGGGC	TGATGTCTGA
38851	TTTTGTCCTG		TCATTTCTGT	ACTACCTTTT	ACAAAACCTC
38901	TCCTATGACC				
			TCCAGCTCCA		CCTGCTGTAT
	ACCCTGTGGG			CTCAATGATG	AAGAAACAGG
39001	CTTGGAAGTT	AAATTATCTA	CCCCAGGCCC	ACAGCCTGGA	ACCTAGGATT
39051	CCAACCAAAC	CTTGTCTGAT	TCTAAAGCAT		CCATACTCTG
			TCAGTTTCTT		CCTCCAATTC
39151	TCACCCAAAC		ACAGTCTCTG		TTGCTCCATC
	CCTTGGCCTT				CAAGGCTTCT
39251			CTTCTTGGAG	TCTCTTTCTC	CCATGTTCTC
39301	CACAACAGAG	CATTCTCCTG	ACTGTTTTCA	TTCTGCATCT	CACTCTTTCA
39351	TCAGTATCTT	TTTCTCTACC	ATGCCCCATA	AATTTGGGTG	CTCCTGAGGG
39401	TCCTGTCCTT			ACAACCTCCT	TGATCTACTT
	CATCTACTCA				ATTCAAATCT
	GCATCTCTAG				
					TACCTGAATA
	TTTTATAGGC				CTCTAAGTCT
39601	AGACTACAGC	AGAAAGCAAT	GCTCTTTTTA	TTAAGGCATA	GTGCCTCTTT
39651	CAGAATAATT	TACAGCATAC	AACCAGGCCT	GCTGTGCAGC	ATTACAATTT
39701	GTCATTAAAA	CTCCATTCCT	CTTGCCAGAG		ATTTACAGCC
39751	AGGGCGCCAA				TATTATGAGT
39801					
		CTCCTCAGAC			TGGAGTTGCC
39851			CCAGGCTCAA		CTGGATATTG
39901		TGCAGAGGAC		CTGAACAGTG	TTCCCCCAAT
39951	GTGGGTGGTG	ATCCTGAGAA	ATATCATTTG	TATCTGCATG	TGCTGTCTCA
40001	CACACACTAG	CTCACATGTG	CACACACACG	TGCATGCACA	GGACAAAACC
40051		CAACCCAGCA		AGCCATCAGC	
	CTTTATAGGG				TCAGGTGAGT
40151					
		TATTAAATCT		AGGAAAGTTA	
40201			GGGCTCAGTA		AAATGTTTAG
40251		TCCTACCGAT	AATCTTTCAG	ATCTCAGAAT	TCCAGCCCCT
40301	TGTGCTGTTC	TGGGTTGTCT	GACACAGACG	AAGCAGAGAA	CAGTAGAATA
40351	AACAGCTCAG	TAAACAATTC	ATTGAGGGAA	AGAGAGTGAG	AAGATTCACT
40401			ACTGCTGGTG		
40451	TAAGGCCTGC		TTGGTCTTAT		GTCCCACCCT
			CTGGCAGGTA		GCGTGAAAAT
40551			CAAAAAATAC	TGGATTCTGC	CCTCCAAGAG
	TTTACTGTTT			CCCTCCTTTC	TCTGCCTCTT
40651	GAAGACTGAC	CTATCTTTCA	AGGCCACTGG	CCCAATTCTG	TTTTCTAAGT
40701	AAGACCACTG	AGTCAGTGGT	GACCTCTCCT	TCTCCCTAAC	AAAGTCTGAT
40751	TTACTTGAAT		CTCCCTCTTG	GCCTGTGAAT	TTCTTGTGTT
40801		TCTGATTTAT		TCCACAGTAC	
40851					CTGGTGTAAA
		AATGCATTGA			AATGCTCTGC
40901			AAGTGCCTTA	AACTTTGTTT	TTCTCTTATG
40951	TAAAATAAGG	ATAATAATAA	TGACACCCCT	ATAGGATTGC	TGCAAGGATT
41001	AAGTGTGATA	ATATATATAA	AACTCTTAGC	ACAAACACCT	GGCTCACAGG
41051	AATAGTAGCT	ACTACCATAA	TGGTAACTTC	GAGGGCAAGT	TTTCTCAGAG
41101	TTATTTAGCC	CTCCTTCACC	CTGTGTCCAG	GAGTGCAGAT	CAGAATGGTC
41151	AGATTCCAGG	ACACCAAGTT	TTCTGTGGGA	CCTTCCCTAC	Charagaacr
41201	A A C C A A TTTTA	A A TO CO COTTO	AGCTCATGCT	COUNTY	COMMINIANCI
41201	TCXCCC* TTC	CCMCMCCCC—	MMCCLATGCT	GITACACTCT	CITCUTCUAC
41701	TCAGGCATTG	GGTGTGGCTT	TTCCAAGCTT	GAGAAGGGTG	TGATCTGAGA
41301	TGGGCTTGGG	TATAGAGGGG	AATTATATTT	AGGTCTACCC	TGTATAGGAA
41351	AAAGTGCCTT	CCCAAAGTCT	CCCTGGCCTA	AAGTATAAGA	GATATGTGTT
41401	GGGATTTAGA	CCCAGAGCCC	AAGCCAATAA	TGGGACCCCC	TTCTCACATG
41451	TGGCTACCTC	CTGCTATCAC	CACAACAGCT	ATCATACCCA	TAACTACAAC
41501	AGAGGCCAAT	TAACGTGGTG	ATAATTGACA	AATGTCAAGA	CATCCTACAT
41551	TCACCCACAC	TOTOCOCOTO	GCGTGAGCTT	MUN N NUMBER	ACCCA ACCA A
41601	2 COMMON A	1010001111	GCGTGAGCTT	TIAAATIGGI	AGGGAAGGAA
41001	AACITITATA	CCTACACCTA	TCATGGAAGG	CAGAAGGTAA	GAGCTAAAAT
41021	AAAGGTATGC	CAAGAACAAA	GGCAGGAAAG	AAGGGTTTTA	ACAACTTGAG
41701	GCCTGATCCA	TTGATTAGTG	AAGAGGAAAC	ATGTTCAAAA	ACCACTCTAT
41751	AACCACCTTC	TCCAAGTTTT	TTATAATTTT	GCTTCTTCGG	ATATCTTCTC
41801	ATCATAGTCT	TAAATGCCAT	CAAATTAACT	GAAAAATGCT	AAAAATGCAA
41851	CCACTCTAAG	AGAATGGGTT	AGATGGGAGA	TCCCTTTCTT	DADCDACTCC
41901	GTCTTAAACC	אאאארתאררר	CTTTGTCATG	CORCORROCC	DOCAROLLO
41051	OTO THUMBO	UNUACTURE CONTRACTOR	CTITGTCATG	GIAGTATGGA	AGGAAGGACA
4722T	CONTRACTOR	MUMUMANAAAA	GTGCAGGGCC	TGTTGAGGAA	GGAATGAGTA
42001	GIAAAATATG	GCTAGAACAG	GGTGCAGAGG	GGAAGAACTT	CAGAGAATGA
42051	CCAAATAAAC	AGGCTGAAAG	GTGTAGACAT	TATAGGCAAT	AAAGCAACCA
42101	CAGAGGTTTC	TAAGCCATAG	GGTGACATGA	TAGATCTGTA	TTCTAGAAAA
42151	GTTAGTTTTG	CAGCAGTTGT	GTCCATTGAA	AGGGACAGGA	TAAGGGAGAT
42201	AGATAAGAAG	ACATGCTATC	ATGATAACTA	CATTTCCATA	CCAACTCCTA
42251	TGGTGGAAAG	GAATGAGAGA	ACAGGGTCAC	DCATCA ATC	CUCCCCARM
42301	TCD A TCC A TC	ATARCACAS	CENTRACE	AGAIGAATGA	CIGCCCAATT
15201	TCAMICCATC	ATAACAGGAT	GTATAGGATT	GCCCTTAAGT	AAGATGGGGA
		~	T ~ T ~		

FIGURE 3, page 11 of 57

42351	ATCCAAAAAC	GAGGAACAAG	TTTGTAAGGT	TTTGGGGGCC	AATGATGAAT
42401	TCCATTTGGG	ACATGTTGCT	TTGGATATAC	CAATGGGACA	TTCATGTGAA
42451	AATGATCTCG	GCAATCCTAT	CCTGGAATTC	AGGATAGGAT	CAGAATGAGG
42501	GACACAGTTT	ATAAGGTAAA	CAGAATGGAG		AGATAAGGGC
42551	ATAGATGAGC	TTACCAAAGG	GGAGAGTTTA	GAATGAAAAG	
42601	AGGCTAAGCC	TGTGCTATTC	TTTCTCCTCA		
42651	CACAAACCAT	CAGTGAGTGT	CATGATAACA	CTACTGTGGG	
42701	CTCTATAAGG	GCCTGATTTC	CTCCTCTATA	AAATAGAGGG	
42751	TGGTCCATAT	CCTGTTAATT	GTGTTTGGAG		AAACCAGCTA
42801	CTATCCAAAG	GGGACATCCC	GAGGCAGGAC		AAATCCAGCA
42851	CAGGGAAAAC	ACTTTCTGGT	GCTGGTCCCA		GTTCAGTTTA
42901	ACCCATCACC	ATCACCATCA	GTAGCTTTCA	GCTGCTACTG	ACCACACTTA
42951	TAGGAAGAAA	AACAATTAGA	ATGGAGAGCT	AACTCTTTGG	
43001	AGAACACGGG	TCTACAAAAC	CGTCAATAAA	GCGCTAAGAT	GCCTGGGCGG
43051	GGTCAAAAAG	TCTACCTGGG	CGGGGTCAAA	AAGTCTACCT	GCTCAGCATA
43101	TGGGGCCCAG	ACATCTGACC	TTTACCAACT	CCACAATAAC	
43151	ATGGATCCAG	TCTTGGTATC	ACCTAGTCGC		
43201	TTTGGTTCTC	AATGGTAGGT	GACTGGAATA		TCTCCCACCC
43251	CTACCGCCAA	TCCTTTCTGC	CCCCTTATAG	TTTAATTTGC	TTGTAAATTA
43301	CTTGGGAATA	CATTTGGGAG	CCATTATAGG	GAAATAGAAG	
43351	TGAACAGAAT		TTTTATTACT	TCACATTGTG	
	AGGAGGAATT	CTAGAAGCCC	CTCCCAGTGG	CCAGGAATTG	GTCATAGCAT
43451	GAATAAACTC	AATATAGGTT	GAGTATTCCT	TACCCAAAAT	GCTTGATACC
43501	AGAAGTGTT:	TTGGATTTTG	GATTTTTTTT	TTGAATATTT	GCATTATATA
	CTTACCAGTT	CAGCATCCCT	AATCCAAAAC	TGAAATCTAA	
43601	TGAACATTTC	CTTTGAGTGT	CATATTGGCA	CTCAAAAGGT	
43651	GAGCATTTTC	AATTTTSGCT	TTT XXXATTA	<b>GGGATACTCA</b>	ACCAGTGGTA
43701	GGTTTGGGAT	GATATCAGCA	TGTTAAGGTC	AAAGAGACCT	AGCTGGGAAG
43751	GGTGGGAGGA	A TATGGAATT	TTCATTCTCT	GGGCACCCCT	TGAACAGTCT
43801	TACTATTAGG	GTCCCAAATT	THITTETAAGT	GTGTGTGTGT	GTGTGTGTGT
43851	GTGTGTGAGA	GAGAGAGAGA	GAGADAGAGA	GAATTTTCTT	TCTTCCTTTA
43901	TATTCTAAGT	TOCTCALGAT	AAAATTTTGG	GTTTCTTTGT	ATTCTCCCTG
43951	CAGCTCCTCA	TGTAGTTCTA	ASCAAATAAA	GGAATTCATT	AGGTCCTTGA
44001	TTTCAGAAGC	CTCCCAUTTC	TETATUTAGE	AGGAATCTTA	GGGTGGCAAG
44051	ATAAGTTGAG	GGACTTTTCT	TEAAGGACAT	TTCACAAGTA	AGAGAAAATG
44101	TTGACTGTGT	ATATCTAAUA	ATHOOTGOOG	CTCAATGATG	CCCCCTAAG
44151	TTACTCTTTA	CTATTATTGA	TIGATIGATI	GATTGATTGA	AGAAGCAATG
44201		TTGAAGAAGT	AATGTTTCCA	ATGGCTACAG	CAGACTGGAG
44251	CAAAAGAACA	AAATGAAASA	AAATACATTA	GGCTTTCCAT	TTCTTCTAAT
44301	TCTGGGGCAT	CTGATGAAGC	TTTGGATCCC	CCAAGGTAAG	AGCTGGACTC
44351	TGCTGGTGAA	AACTCTTTAG	GAAAAACAAA	AGAATATTGT	CAGAATCTGA
44401	TGCACCTTAG	AAATGATGCA	GCAGAACTGC	TTTATTTTCT	AAAAGGTGAA
44451	ATGGAGACCC	AGAGAAGCAA	AGTGATTTGT	TCATGATCAT	ACAGCTATTC
44501	AGTAAAGCCA	GGACTTCTGT	GATCCACTGT	CCTTTCCTTA	AACCAGTGGT
44551	TCTCAACCTT	GGGAGCTTTA	AAAAACTGCT	AGTGTTGGAT	CCATCTCAGA
44601	CTAATTAAAT	CAGAACCCAT	GGGGATGAGG	CCCAGACATG	AGTGGGTTTT
44651	TTGTTCTTTT	AAAAAAATT	GCTCCCTAGG	AGATTTCTCA	AAGAACTGAA
44701	AATAGAACTA	CCATATGATC	CAGCAATCCC	ACTTTTGGGT	ATCTACCCAA
44751	AGGAAGATAA	TATATTATTA	AAAAAAGATA		AATATTTATT
44801	GCAACACTAT	CCACAGTAGC	AAAAATATGG	AATCAACCTA	ACTGTCCATC
44851	CATGGATGAC	TGGATAAAGA	AAATGTGTAT	ATATACACAC	ACAATGGAAT
44901	ACTATTCATT	CGTAAAAAAG	AACAAAGTCT	GTCTTTTGCA	GCAATATGGA
44951	AGGAACTGGA	AGCCATTCTC	TTAAGTGAAG	CAACTCAGAA	ACAGAAAGGC
45001	AAATTCCACA	TGTTCTCACT	TACAATTGGG	AGCTAAATAA	TGCATATGCA
45051	TGGGCACAGA	GTGTGGAATA	ATAGACATTG	GAGACTCGGA	AGGGTGGGGG
45101	GAATGGGAGA	GGGTCAATGA	TGAAAAATTA	CTTAATGAGT	ACAACGTACA
45151	TTATTTGGGT	GATGAATACA	CTAAAAGCCC	ACACTTTACC	ACTATGCAAT
45201	ATGGCCATGT	AACAAAATTG	CCCTTACACC	CCTTAAATTT	ATACAAATAA
45251	AATAAATAA	ATAAAAGCTC	CTTAGGGCTG	AGAACTACTG	CTCCTGTCCT
45301	ATGGGTCCCC	AGCTTTATTT	TAACTCAAAA	TGAGTTTAGA	AAAATTTATG
45351	AACCCATTTA	ATTTATAAAA	TTGAGTATCT	CCTGTGTGCA	AGGCACTGTG
45401	TTATGTTAAG	TGGCTGAAGG	GAAATTAGAC	TGGGGAAAAA	GACAAGGTCA
45451	TGGCCTAGGT	TTCAAACTAA	TATAAAAGAC	ATAACAAATA	AGAAAGGATG
45501	CCACCTTCTT	CCAACCCTCA	TCCCTCTTCC	TTTTGACAGT	TGCAGATGTT
45551	GCTAATTCAT	TTTGGCACCC	TTTTTCTCTG	ACCCAAATAT	AGTCTTATAA
45601	ACCTTTTCAA	CCCACGGCTC	TAGGCAAGTA	TCACCTTTTG	CTCTTTTGGC
45651	ACCAGATCTC	TTGAACACTA	TTTACTGGTT	TTGGAAAGAT	TATACATGTA
45701	TGTCTGGAGT	TGAATGACTG	AACAGAGCAA	TAATAAGAGT	TAAAGCAAGA
45751	AAGACAGGCC	TACAGGAGAT	GGCAGAGGGT	CTTGCCTGTC	AGGCATTGAT
45801	TTTGAACTTC	ATTGCATAGG	CAATCAAGAA	CTATTGAAGT	TTTTGCACAA
45851	AAGACTATAG	ATGAGATTAA	CCTGGTTACC	GTAAAGGACA	AAGTGATTGC
45901	AGGTAGAATG	AGGCCAGCTT	CATAAATGAA	TCATCAGGAT	ATGAGAAGCA
45951	AGGGCTTGAA	CATGAGAGGC	CATAGTGGGA	ATGGAGGGAA	AGGGACAATG
46001	TGAGAAGCAG	TGAAGGAGAA	GGGCTGATTG	AGTAAAGCAG	TGGAGAAGAC
46051	AGTGAAAGAT	GTCAGATGAC	TACCATGTTT	GGCGACTGAG	TGAGGGAAGA
46101	GGTGGTGATG	ATATTACTGA	AGAGAGAGGC	AAGGGGTGGT	CACTGGATTT
46151	AGAGCAGACA	TTATCAACTT	GTGGTGTCCA	GACATTTCAC	CCTGGGAGAA
			YOTTO		

FIGURE 3, page 12 of 57

		AAGTGGCTTC			
46251		ACTGACTGAA			
46301		GTCTCTTCTG			
	ATCACAGGGC		ATTAAGTAAG		
	TAAACTGTAA		ACAATTGGAT		
		GATGTCTGCT			
46501	GCCAGACCTG	GAAATGGAAA	CTTGAAAATG	TTCTCTGTAG	AAAAGATAAT
46551	TAACATTTGA	GGATGGTTAA	GTCCTCTTAA	<b>ATAGATGTCA</b>	GAAAAAATGG
46601	AGGTCATGTA	GACAGAATGT	TGGATAACAC	TACTTTGTAA	AATATTTTAT
46651	CTTATTTCCA	TTATAAAAGA	AAAAAAGCTG	GGCTGGGCAC	GGTGGCTCAC
	GCCTGTAATC		GGGAGACTGA		
46751	TCGGGAGTTC		TGGCCAACGT		
46801	GAAAATAGAA	AAATTAGCCG			
46851		GCTGAGGCCG			
	TGCAGTGAGC		CCATTGCACT		
		CTCAGAAAAA			
		TAGTCTACTA			
	TTTATTCCAT		TTTAATATAA		
		GAGAGAGTGA			
		AGAGTATTTA			
		CAGGGTGGAG			
		CGGGTTCAAG			
47301		GGCAACCGCC			
47351		GATGGAGTCT			
	GCATGATCTT		AAGCTCCGCC		
		CCTCCCAAGT			
	CCAGCTAATT		TAGTAGAGAC		
	AGGCTGGTCT		GCCTCAAGCA		
		GAATTACAGG			
47651		ACAAGTGAGA			
47701		GTCATCTACT			
47751	TIGTIGTIIG		CTTAGTTTAC		
		CTTGGCTAAG			
47851	CCCAGCAATC	CTAGGAGGTA	GGGTTTATTA	CTACCCATCG	TACAGAGGAG
		CATAGAGTTT			
47951	AAGTGGCAGA	GCAGGCCAGG	CCAAGTCTGT	CTGACATCAG	AGCTCATCAG
48001	AGCCCTCCCC	ATTGTCCTTG	AACCAGTAAA	GATGGAGTTC	TTCTACAGGG
48051	GTGGTTGGGG	GACAAGGACC	CCATGGGTGT	<b>GTCTGAGTCA</b>	GAAACATCTG
48101	CGAGTGGGCT	GAGAAATGAG	TCTTCTGTGA	AAAAGAGCAA	AAGAAAAAAT
48151	GGGTCAGGAG	CCAATAATCA	TTGTCCATCT	TTGTGTGAAT	GTATGGTGTG
48201	GGAGTGGGAG	CAATAAACGA	TTCTAAGGTC	ACACAGAAAA	GATGCCACCT
48251	TCTCCAATCA	CATACCGCCC	CTCGTCCCCC	AGTTTTCTCT	GAAATAGCTC
48301		TCTATCCTGG			
48351		GACAGGGATA			
48401		GAGTCTGAAT			
48451	TGGAGTGAAT	ATTGGGCCAT	GGCCCTTTTT	CTTGCCAGCT	GAGCTATGAA
		CCTAAGACCA			
		GAGCTGACAG		TTGTTTTTT	
48601		GTAAAATTTT			
48651		CTGGGCTCCC			
48701		TTCCCATGGC			
48751		TCCTTTCAAC			
		AAACCCCCTA			
		TTTTTATGCC			
		TGGTTTAAGG			
48951	CACATGTCAA	AGCTGTGGAC	GTTAGACTTG	DCCDCDCCAA	CACATATCAC
		AGGAGCATCT			
		GTTTTTGAAG			
		GGCTCCTCCA			
					GTAGGGAGCC -
		CTCTTCAAGG			
		GAGGGGTTAT			
		ATTTTTTGAA			
		GTTGCAGATT			
		GCTTTGTGAA			
49451	ACATCAAGTT	AAAATAGAGG	AAGATGCCTA	GAAATGGCCG	TATAGACAGA
49501	GAAAACTGCA	CTAAAACTCC	CTCCGTCATG	CCTGACTCCT	CTCTAGACTA
49551	TGACCATCGA	GGGGCCAGAA	ATCATATCTT	AAAGATCACT	GTGCCTCCAG
49601	TACCCAGCAC	GGTGTTTAAT	AAATGTTTGT	TGAATGAACG	AACTAGTAAA
49651	ATTTTCAAAT	CATTAGAGCT	GAAGTATCCT	TTAAGATTCT	TTAGTCCCTC
49701	ATTTTACAGA	TAAGGAAGCT	AAGGCTCAAG	ACATTGTGTG	GCTTGGCCAA
		CAAGCTAAAG			
		TGCTACACTT			
		TTTTACAGGC			
		CTCATCAGAA			
		CTCTGCCTCT			
50001	AGTGGTTGTG	TATGATGATT	AGTGTAAGTA	GGATGGGCAA	ATGCACACCT

FIGURE 3, page 13 of 57

:	50051	TTCCCACCTT	CAAACTCAGA	AGTTGTAACC	AAGAGTCACA	CTGACTAAAC
:	50101	ACTCCAATTT	CCCTTTCTGT	TTTTCTTAAC	ATATGTCCTA	
:	50151	AATAGCCATG	GTATATTAGT	CATGGTATTT	CACGCTAGCT	
	50201	CTTCCAAATC	TCATTGGCTT			
-	50251	AAAGTTGAAT		GGCAGTTATC		
	50301		CACCTTCCAT	CGTATTGGCT	TAAGCCACAA	CTTGGGGATG
	50351	ACTTCCTCTC	CCAMAAMCCA	CGTATTGGCT	CCACCATTCA	GGGATGGCAG
		CACCOUNTE	GCATAATCCA	ACCAATAGAG	GGGGGAGGTT	TGGCACTTGT
	50401		CCTAGCCTAG	CATTGACACA	CACCACTTCT	ACATACACTC
	50451	CCCTAGTCAT	CATTCAGTCA	TGTGGCCCAA	CCTAGATGCA	AAGGCATCTG
	50501			GTCAGCAACA	ACTTTGCACT	TGGAAGGGGA
5	50551	GCCTGAATCG		CCAACACATG	TAACTAGCAA	
5	50601	CGTTATTTGT	CAGGCAATGT	GCCAAGAATT		ATCTTCACAA
5	50651		AGGTTATTGT		GTATAGATGA	
5	0701	<b>GGTAGAGATA</b>	TAACTTAACT		TGCATAAGTG	
5	50751	AATCCAAAAT		CTCTCCAGAG	CTCAGGCTCA	
5	0801	ATTCTACTGC	TTTGAGCTTC			TGATTGCTGC
	0851		GTCTCCAGCA			GCCACTAAGT
	0901	ACCCIGIGIA	DICTCCAGCA	ATTACTTTCC	TCCCTCTGGA	TCTTGGTTTC
		CCACCECCA	AAGTGAGGAT	GTTTAACTGG	ATAAAATCTG	ATGTCACCTG
	0951		CATCATATGA		AAGCATATCA	
	1001	GTCCCCAGTG	ATGCTTGACC	ATAGCAAAGC	CCTTTCAAAG	GTTTCTTAGC
	1051	ACACCACATA	AATGGAAGCC	TCACAGTGTC	CATGTAGGAG	AAAGCAGGGC
5	51101	AAAGTATTTC	CATTTACCCA	ACAAAGAAAT	CAACATATAG	TAAAAAGAGA
5	1151	GTGTTTTCCC	ACCAAGGCCT	CAGATTGACT	AGCGGTAGCC	TTGGAAATAG
5	1201	GACTTTATTT	TGTATAGTAC	TTTTGCCACC	AGGGTGGGG	GGAAAAGAGT
5	1251	GCTTCTTTGC	CCCAAATGCT		AACCTAAAGA	TGTCACATGG
5	1301		TTCCCCCAAT	CCCCTCAAA	ANACTACTEC	CACTTAAATG
	1351	AAAGAGTAAA	GCTGTAGGAC	TTTTACTCACC		
	1401	CACTCCCATC	CTCTTGAGGG	COMOCAGO	AGTGTCCTGT	GGGGTCCTTG
	1451	TTCTCCCTTAC	ACCERTED AS	GCTCGAGGTG	TATGAATTCC	CCAGCATTAC
	1501		AGGTTTCAGA		GAGCTCCAAA	
		ACCCAAGTAT		AACAATCCTT	GAATGACTTT	ATACAGCAAA
	1551		ACTGTGTCCT		TTGTGTGTGT	TTGTGTGTGT
	1601	GTGTACAACT		TTTCTACCTA	TGTCCCCCTG	ATGCCTCCAC
	1651	ACAGAACATC	CCAAACTCCA		CTCTTGAGAT	TCCCAAACTT
5	1701	GGAAACAGGA	GATGCTTCAA	AGGCCTCTTG	GAATGTCTTT	TGAGGCTTTA
5	1751	TATTGTGATA	TGTTGGACAG	ATGGTTAAGA	AACAGAAGAA	GAGCATCACC
5	1801	AAAAGGATTT		GTGGAGATCT	ATTAATATTT	GCCACTAGCA
5	1851		TTCTTGGGAA		CCCTAGAATC	AGATTGACCC
5	1901		AGGGAGAATA	AATAGAGACT	TGAGCTTAGA	
	1951	TGGCCAGAGC	TGAAAAGGCT			
	2001	CTTCAGAAGA			CAGAGAAGAT	GCAAGAGCAG
	2051			TATTTGGGTA	GGTTCCTCTG	GTGTAAAGGG
-	2101	CCCTCCATCC	DARCECCOR.	CAGAATAAGA		
		CENCOMICO	AAACAGGGTA	TAAAGCAGGA	GCATTTGGAA	TCTGCCCTTT
	2151		CCAGAGAGCG	TCAGGCAGCT	TGTTGGGTAA	TAAGTAACAC
	2201	TGGCATTTTT	CCCATGGTTC	TGTCATCTTA	AAGAGCAGGA	TACATAAAGG
	2251	GATTCAGATG	TCTTGTTGGT		TTCTTTTTAA	TACCTTGTTT
	2301				GTGTGGTAAG	ATGCACAGAC
5	2351	ATGGAGATGA	CAGTCATGAA	GGAAGAAGTA	TTTATACTCA	
5	2401	TAAATAGGAA	GCATGGCCTC			CACCAGGGTC
5	2451		CAGAAGGAGC		CATGGACAAG	AGGCTCTACT
5	2501		TGGCAAAGAA		AGAGTAAGCA	GGTTTAGGAT
5	2551		GAATGACTTG		AGGGTGTAGA	
	2601		ACCAGGGGTA			GACTGCCTCT
			CTARACCACC	MITAGGGCAG	CTGGATAGTG	GTCTGGAGTG
5	2701	ATCTCTANAM	CIAAAGGAGG	TGGTTGGAGG	TGTAGGTTTT	GGATTGGTTG
5	2761	ATCIGIATAL	GMANGGIGCA	CGTGCAGGTT	GAGTCCTCTA	CTATCACTAG
2	2001	AMAITGGCIG	GTCCCAGGAG	AAGTAGTCTC	TCTAGAGACA	GCAATGCCCC
2	2001	AGATGTCAAA	GCATCAGAAA	ATACAGAAAA	AAAATTAAAA	GCATGATTAA
5	2851	TTCATACTCA	CAGGTCTAGT	TTTTGTGTAG	TTAAGAGCAA	CCTAAAGAAG
5	2901	TTGATAACTC	GTGTTGCAGG	TCAGGTTTCC	CAGAAATCAT	ATTCTCAGAT
כ	2951	GAAGATTTGC	ATGAAGGAGG	TTTAATGCTC	AAACTAAGCC	CTAAGGCTCC
5	3001	ATACCTGTGG	AGGAAGTGAA	AGAAGCCCAA	CTGGGCACAG	AAGGTGGAAC
כ	3051	ACAATGCCAG	TCACACAAAG	ACCTCAGTGG	ATCCTGGGCC	ATGAGGAGCT
5	3101	CTAAGCACAG	ATGACCCTTC	AGAAATGTCT	CCAAGTGGGG	AAAGGAATCA
5	3151	TGCTAGTCAC	TGGATGTGGG	CTTCCCACTC	CACCCCATGA	GGCCATCACC
5	3201	TTAAGTGAGA	GAGCTCTTTG	GACACAGGGC	ATCCTAACAC	CCCCACMONA
5	3251	CACCCACATT	GCCCACCAAC	ACTICACA CCA	ATCCTAAGAG	GGGCACTCAG
5	3301	TCCCAAACCC	CARTCECCA	ACTCTCAGCA	GCTAGAAGAA	GAAGGTATAG
_	3361	CTACCCTCA	GAMICIGGGC	TGCACACCTT	AGTATCCATT	AGAACTGGAA
2	240-	GIAGGCTGAA	TUCCAGGCAG	GGATCCCCTG	GAGAACACAG	GTAATTTTTT
5	3401	MAAAAATCAA	GCTATGTGTC	TGAGGCTATG	TGGTAAGACA	TCTCAGTTTT
5	3451	CTGCTAGGAA	AAGCCACCAA	ACCAGATTGG	CTTATTCATG	TTGAAAAGTC
5	3501	TGAGAATCAC	ACTCAGATGT	TGTTGATAAT	TCTGCTTGGA	ΤΑΔΔΔπητηΔη
5	3551	CTATTGGTAT	GCTTGTGATA	TAGCAGTACC	ATTGCTAAAA	ATTCCATGCG
כ	3601	GAGAATCCAA	TCTGCATCAT	TTTCTTTCTC	AATGATTTGT	TTTTDDDGGC
כ	3651	AGAGGTTCGG	CTGTGCCCCT	TTAAACCTTC	TGTGCAAGTG	CCACCTTCCT
5	3701	TTCAAATGGA	GAAGCAGCAG	CCCTGTCAGA	ANGGGTGGCT	GGAGCTCCC
5	3751	TTTTGTGAGA	GGAGGAAAAC	TTACTGGGAA	TTACCTTGGC1	CACACCCCCC
5	3801	CATGAAGGCA	TACCACTGCT	TCCTCTGACC	**************************************	TAGAGCCACA
5	3851	ACATACTOTO	CTACCTOCI	ACCAATGATG	1 TCCAGCCGG	TATATTAATG
-			O. ACCIGAGA	ACCMATGATG	MAGTGGGTGA	TGTGCCTGGC

FIGURE 3, page 14 of 57

53901				GGAGATGATA	
53951				CACCATGTTC	
54001	AAGTTCTCAA				TTCCCGTTCC
				CACACACACA	
54101	CACACACAGT			AGTTAGTATG	
54151				CAGACAATTA AAGCGCCATA	
54201	TGGGAGCGGG			AGTTCAAAGT	
54251 54301	CTCTCTGAGC			GTGGGGGAGG	
54351	TTCTCTTTCT			AAATAAAAGC	
54401				GCGTCTGACA	
54451				GATACTTTTT	
	AGGACCTCCG	CTTCTTCCCT	TCTCACATGA	GAAGGAAGAT	TTTTCTAGAA
54551	ATCTACAGGT	GTTTAAGCTG	GAATGTGCCT	CAGACATCAT	CTGGTTGGAC
54601	CCTTTCATTT	TGCAGATCTG	AGGCCTAGAA	AGATTTGGTA	ACTTGCCCCA
54651	GGTCACAGTT	GACAGAATTG	CTCAGTGAAA	AGTCCAGCAT	AAATACCCCA
				TAGTGAGGCA	
54751				CTTAGGACAG	
54801	CTCCTACTGT			TCAGGCACAG	
		CCTGATATGT		ACTGTGTGCT TCTTTATAAT	
54901				CGAGGCAATG	
	CAAAGAATTT			ATTTTGCACA	
	AATGAAAAGG				AATTTCTTTA
				TCAAATTCTG	
55151				GCTCGGGCAG	
55201	TCGAGTGCCA	AGTGCGCTGC	GCTGGGGTCT	AGTCCTGACT	CAGCCGCCAT
55251		GCTTCCTGGA			CTCAGCTCCT
55301	TCATCTTAGG	AAAGAAGGGT	AAAGATCTAC	AGACAAATTG	ATCTTTAAGT
	ATCCTTAGAG				ATCCTTCCAA
55401		GTAGGGAATT			AGGGAGCCGG
55451		GTCAGCCCCA			TGTGTATCTG
55501				AGGAAACACC	ACCGCTCCGA
				CACACCCGCT	CAGGAGAGGA
55601				CTCTGCAGAC	TCCCAGCAGG
55651				ATGCTGTAAA	
55701	TCTTGCTTGG			AAGGGTTATA TCAAAGCTAT	TAATTGCAGA TTTTAATGGA
55751			CAATTTAAAG	GGAGGAGAGA	
55801 55851				AGCCATTGCA	
55901				CAATTAAATC	
				CTACAGTTTT	ATTTTGAATA
56001				TGAATAAGAG	ATCTCAGTCT
56051		CAGCCAAAAG	CCCTTAGTGT	GTCCTTGATC	<b>AAGTTACTTC</b>
56101	CCCTATCCAT	TTCCTTACCT	GCAAATGAGA	AGCTTGAACC	AAACTATCCT
56151	AATGTCCCTT	TCAACTCTAA	AATCCTAGAT	GATCCTCAGA	
56201				GGTCCGCAGA	
				GACTCCTCAG	
56301	ACACAACCTT			AATAACCTGG	
56351	TACAAATTTT	CTTGGGGAAG	AAGGTAAAAG	GGATCTAGCT	TTCTGGGTTA
56401	TGAATGCCAT	GTAGGGAGGG	CATGGTTTGA	GTTAGTCCTG	CCTCAACACC
56451	. TTCATGAGAC	TTATICICAA	ATCTTCAGAG	AAGAAAATTC	TCACCTTACC
56561	CTCCTCCTTC	**************************************	TTGAACTTCC	TATAATTGAA	CAAGATAAGC
56601	ATCTCACCTA	. IGHIIIGACC	TTABABACTTO	TAGCTTCCTT	CAGCACAGAA
56651	GTGGCTCTCT	GAACCAATTT	TAAGCAATCO	TGGCTCTATC	TGTGCATGTT
56701	GATTTAGCCT	GTGGTTATAG	TGTTAACAAT	TTAGTGATTC	ACCTCATTTT
56751	TAATCTCTCT	TTCCCTTTAC	CAGGATCATT	TTCTCTGTGT	TAAGGGATCA
56801	ACATTGAGGT	AAGAATGGCT	' AAATAATAGO	: ATCTTCTGGA	ATACAAATGA
56853	L CTTTATAAA1	' AAAAGAAGAT	' AAAAGGAAGA	AGTAGGATGA	TTTCTCAGCT
5690	L CTAATACACT	' TAGCAAATGO	CATATGCTT	CTCCTGCGTG	TACTGGTCAG
					TGGTCAACAG
5700	L TGGATTGCAT	ATGTGACGGT	AGTCCTTTAL	A GATTATAATA	CCATATTTT
5705	I GCTGTGCCTT	TTCTAGGTCT	AGATATGTT	r AGATACACAC	ATACTTACCA
5710	1 TTGTGTTCCA	ATTGCCTACA	A GTTTCCAGT	A CAGTAACCTG	TTGTACAGGT
5715	1 TTGTAACCTA	GGAGCAATAC	GCTATACCA	PACAGCCTAGO	TGTGTAGTAG
5720	L GCTATACCAC	TTAGGTCTGC	STAAGTACA	TCTATGATGT	TTTCACAGTG
5725	L AIGAAACTIC	. CIMMIGHUMA COTACTATA	T CTCDDCDCT	2 #CCCCDDGC0	GAAGTCTCCA
5736	L GAGGCAIGAC	AGIACINIA AGIACINIA	P DTTTAGGGGGGGG	ATCGTGGAC	CCAAAAGTTC
5740	I DESTRUCTION	ACADCHAIG	ב הבינהטונט	T CARGTCCAC	CAGAACCTCA
5745	I GAATACAAT	TTATTTAGA	ATAGGGTCT	T TGCAAATGT	GTAAGTTAAG
5750	1 ATGAGGTCA	CACAGAGTA	AGTGGGCCC	T AAATCCAATA	TGACTAGCAT
5755	1 CCTTGTAAG	AAAGGAAAA	GAACACAGA	CAGGGGAGAA	GCCATGTGAG
5760	1 AACAGAGACA	A AAGACTGGA	S TGAGGCATC	T ACAAGACAG	GAACACCAAG
5765	1 GATTGCCAG	S AGCCACCAG	A AGCTAGGAA	G AAGCAAGGAI	A GCATCCTCTT
5770	1 CTGGGGCCT	r CAGAGACAG	G ATGGCCCTG	C TGACACCAT	r gtttcaaatg
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FIGURE 3, page 15 of 57

57751	TTTAGCCTTC	AGAACTGTGA	GACAATAAAT	GTATATTGTT	TCAAACCATC
57801	CAGTTGGTGG	TACTTTGTTA	TAGGAAACTA	ATACATTCAG	GATGGAGAGG
57851		GCCCATGAGA	ACAAATGGAA	AGAGCCAGAA	GCCCTCAACC
57901				CCCGCATCCA	
57951 58001		CTGTGAAACC		CTCCACCGCA	
58051	TCACTGACTG	CCTAGACTGT	CAGGGTTGTG	TGGCAAGGCT	TTCATGCCTC
58101	CCTTTCTCAT		CCCCTCAATG	ACTGGTCCAC	
58151	GTGATTTATT			CAGTGCCTCG	TCCTCAAGAG
58201	GGAAGGCATG			CTGTCCTTCC	
58251	TTCTCTGGCA		CCTTGCACCA	CTGTGCTCTA	GATCCTGACC
58301		GTCTCTCCCT	CTCCCTTTCC	ACTCTCCCTT	
58351	GTATCTTTCT	ACAAGCAGGT	CTTCCAAAGT	ACTGCTTGAG	GTCTGAGTTG
58401	GAGGGAACAT	GCCTCTACCC	TACTAAAAAG	AGAAATTCCT	CTGCAGAAGA
58451	CCCAAGCTGA	CTGACAAATC	CCTTTACTGC	AACTGCAGCT	CTAGCTCCCA
58501	CCATTTTCCT	GTACTTACTC	TCCTGCTCAG	GTTCCCTGGC	ATTGCTGATG
58551	TCTTTCAGCC	TTTGTGCCCT	GGCCCCTTTC	CTCCTCTCCC	CTCATCTAGC
58601	ACTACCTGTC	AAAATCAGGG	ACTTACTTTA	AAATTTATCC	CAAATTATCA
58651 58701	TTGCCATCAT		ACCTTATCAT	ATGTTTGAAT	
58751	TTTCCCAAAT AGAGCTGAGA	GTTTTCGCAT		AATTGAGCCT	
58801	GACATCACAG	AAAGTAAGTT	TTTATGAGTC	CTTTGACAAA	
58851	TTCTTTGTAT	TTTTATTATC	GCCAGCCGAT TCCCATTATA	ATGTCACTGT	
58901	ATTTCTACAT	TGGTCATATC	TTTCCTTCTG	TACTCATCT	TGTAATGATT
58951	AACAAATAAT		CTTTGCATCA	CACTTCATCC	CGCTTATGAT TTTACAAAAT
59001	GCTTCAAATT	CAACATGGCC	CTTGATCCTG	AAGATATTTA	TCACTTAAGA
59051	ATCATTATCG	CCATTTTAAA	ATACAAATTT	ATTACTTGGG	CTAAATTTTC
59101	TTATTATAGT	TGGGATAGGC	CTTCATCCAT	AGGGTGAGTG	CAGTATTTGT
59151	GGACTGTCAT	GGCAGCTTAA	ACATTTAGTA	CTTGAAAATC	TGATGCATTG
59201	ATCATCAGAG	AAATGCAAAT	CAAAACTACA	ATGAGATATT	ATTTCACCCC
59251	AGTTAAAATG	GCTTTTAGCC	AAAAGACAGG		TGCTGACGAG
59301 59351	GGTGTGAAGA	AAACGGAGCT	TTCATACACT		ATGTAAATTA
59401	GTACAACCAC TGAGCTACCG	CAGGGAAAAC	AGTTTGGAGG		AACTAAAAAT
59451		TGTGATCCAC	CAATCCCACT AGAGGTATCT	GCTGGGTATG	TACCCAAAAG
59501		TCCCAGCACT			GCGGTGGCTC
59551	TCAGGAGATC	GAGACCATCT		GAGGCAGGCA	GATCATGAGG
59601	AAAAATACAA		AGGCGCGGTG	GCGGGCACCT	GTATTTCCAG
59651	CTACTCGGAA	GGCTGAGGCA		ATGAACCTGG	GAGGCGTAAC
59701	TTTCAGTGAG	CCGAGATAGC	ACCACTGCAG	TCTGGCCTGG	
59751	GAGACTCTGT	CTCAAAAAAA	AAAAAAAAA	AAAGAAAGAG	GTATCTGCAC
59801	TCTCATGTTT	GCAGCAGCAC	TGTTCACAAT		TGGAAGCAAC
59851	CTAAGTGCCC	ATCAACAGAT		AGAAAATGTG	GTACATATAT
59901 59951	ACAATGGAGT	ACTATTCAAT	AAAAAAAAG	AATGAGATCC	
60001	AACAACATGG CACAGAAAGA		AGATCATTGT	GTTAAGTGAA	ATAAGCCAGG
60051	GCAAAACAGT	AAAACATCTT TGAACCTATG	ATGTTCTTAC		GATCTAAAAA
		GGTGGTGGGG	GACATAGAGA GGCTTAGGGG		GGTTACCAGA
60151	GTACAAAAAC		AATAAGGCCT		TGGTTAACTG
60201	GGTGACTATA		ACGTAGCTGT	ACTATTTGAT ACATTTTTAA	AGCACATCAG AAAACTTGAG
60251	TATAACTAAA	TTGTTTGCAA		AAATGCTTGA	
60301	ATGCCATTAT	TCATGATGTG	CTTATTTCAC	ATTGCATGCC	TCTCTCAAAA
60351	CATCATATGT	ACCCAATAAA	TATATACAAC	TACTACATAC	CCACAAAAAA
<b>60401</b>	TAAAAGTAAA	AAAAAAAATT	AAGAAAATAA	AACAACAAAA	CTACATCTAT
00451	TCTACATGTC	TCCATATTGT	AAAACTAGAA	CCAGTCAGTT	AACTTTACAC
POSOT	GAAGGGGATT	GTGGACTTGA	TATAAAGACA	ΔΓΤΤΤΔΤΔΔΤ	ATCCACACCA
60601	GCCTAATCCT	ACAATTGTCA	AAAAGTATAG	TGGATTCTTT	ATTTATTTGT
60651	CTACCTACCA	ATAGAGGTCA	TTTCTGCTTT	AACAAGTAGG	TGGGAGATAG
60701	TEGETEGACA	TECTCCCTCA	TCTTATTTTT	TATTTTAAAA	TATTGGGCTG
60751	AGGCAGGCAG	ATCACCTCAC	AACCTGTAAT GTTAGGACTT	CTCAGCACTT	TGGGAGGCTG
60801	TATGGTGAAA	CCCCATCCCT	ACCAAAAATA	CAAAAAAMMAC	GCTTGGCCAA
60851	TGGCATGCAC	TGTAGTCTCA	GCTCCTTGGG	AGGCTGAGGC	ACCACAAAAA
60901	CTTGAACATA	GGAGGTGGAG	GTTGCAGTGA	ACTGAGATTA	CCCACTCCA
60321	CTCCAGACTG	GGAAACAGAG	TGAGACTCTG	TTTTATATAT	ΑΤΑΤΑΤΑΤΑΤΑΤ
PIOOI	ACACACACGT	ACATATACAT	GTATATATAT	ACACATTATT	ATTCABACCA
PIOSI	GCCAAAGAAA	AATAACACAT	TATATATAGA	GAAAGAGCAA	ATCATCACTC
P1101	ACTTTATATG	TATATATATG	TGTGTGTGTA	ΤΑΤΑΤΑΤΑΤ	<b>GTGTATATAT</b>
PTT21	ATACATATAT	ATATATAGGT	TAAGAACCTT	CAGCACATGT	<b>ልጥል</b> ርርጥልምርም
<b>61501</b>	AACAAACCTG	CATGTTCAGC	ACATGTATCC	<b>CACAACTTAA</b>	ACTENDANAN
61301 01521	AAAAAAAAAGA	ACCTTCTGCA	TGCCAGTAAC	TGTGCTAAGT	GATTAGGATG
61351	ACCCANANCE	MAAAACAAAG	TCCCTCTCCT	TAAAGAATTT	TCTATTTAGA
61401	AGTGCTACAC	ATCANACACA	AAATAAATAT	ATAAATTACA	ATTTGTGAAA
61451	GATTGAGAAG	GGCTCTGACA	GCTGAGACAG AAGCAACATT	ACATCAATGG	ATAAACTTTA
61501	TAGAAGTTAA	ACAGGCAGAT	ATTGGTGAAA	CACCACTCCAA	GCCACACCC
61551	ACATCATTTG	CAAAGGCCCA	GGGTAAAGAA	GATCCTGCTA	AGGADATCAC
					CALL

FIGURE 3, page 16 of 57

			GCAGGACTGT		
			TGCAATAGGA		
			ATGATTGCTA		
61801			AAACTTGTGT TCAGAGCAGT		
			AAAGGGGCAT		
			CCTGAGTCTA		
			AGGGACATTG		
			TTTTTTCTTT		
62051	TTTTTGAGAC	GGAGTCTTGC	TTTGTCGCCC	AGGCTGGAGT	GCAGTGGCGT
62101	AATCTCAGCT	CAGTGCAACC	TCCGGCTCCC	GGGTTAAAGC	GATTCTCCTG
			GGGACTACAG		
			AATGAGTCTG		
			GAGAATCTCA		
			TATTGAGACT GTGAATGAGA		
			GACATGAGGG		
			CCAAGAGAAG		
			ATCAGAGAGA		
62551	CTGAATGATG	AAAGGAGGTT	TTTGGAAAGG	AAAATAGAAG	GGAAGGACAA
			ATATTTATCT		
			AGCCCCAAAT		
			GTATTAAAAA		
			CCCTGTCTTT		
			ATTCGCATGA		
			TTCTGTTTGA		
			CCCTACAGTG		
			ATATATAGAT		
63051	TGGCTGCTGG	CAACATTTAT	TACAATCTGA	ATGTGAAATG	GCTATTCTGT
			TATCAGCCAC		
			CAGTTATCTG		
			AGAGATTTCC		
			ATTTGGTTTT TGTTGAAATC		
			CAGCCTCAAC		
			TGTGGGATCC		
			CTCTGATGGG		
63501	ATGATTCAGC	TGAAGTTTCA	ATGCTTCTGA	AATTTTTTCC	TGCTCCTGCT
63551	GGAGAGCTTG	TTTCTTCTGG	ATTCCCATAG	GTCAGGTCCT	GTGTTTGGCA
63601			ACATAGCATC		
			AGTAAATACA		
			AAATAAAGAA		
			TCTTCTTCCA ATACTGGCAC		
			GATTATTACC		
			CATAAAATAA		
			CATTGTATGA		
64001	ATTCTTCACT	TTCCAAAGGT	GCCTCTAAAT	ACTAAGATTT	CAGTTACAAT
64051			AGATATTAAA		
			TAAAAAAAGT		
			TATTTATCCA		
			ACTGCACTGG ACATTCTTGC		
			TCATAGATGC		
			TATGTTATTC		
			TTTGAATTTT		
64451	TTGAAATGAT	CATATCGTTT	TGCTTTCTAA	AGCTTCTAAT	ATGGTTTAAT
			ATGTGAAGCA		
			TGTTATCCTT		
			GTATTTGTGG		
			TTGTAAGGTT AAAGTAAGTC		
			AACGTTGAAA		
			ACTATTTGGA		
			AATTTCTTTA		
64901	CTATCTTACC	TTGTGACAAT	TCTGATGAAT	TGTGTTTTTC	AAGAAGTTTG
			AAACTTACTA		
			TGTCTATAAG		
			CTTGCTAGAG		
			CTTATTTGAG		
			TCGATCTCGG TGCCTCAGCC		
			GGCTAATTTT		
65301	GGGTTTCACC	ACGTTAGCCA	GGATGGTCTC	GATCTCCTGA	CCTTGTGATC
65351	CACCTGCCTC	GGCCTCCCAA	AGTGCTGGGA	TTACAGGCGT	GAGCCACCGC
65401	GCCTGGCCGA	GGTTTACCAA	GTTTATTAAT	CTTTTCAAAG	GACTACATTT
			· · · · · · · ·		

FIGURE 3, page 17 of 57

65451	TGGCTTTGAT	AATTTTTCCT	ATTTTTTATC	TACATTATAC	TGATTCCAAT
65501	TCTTATCTTT	ATTCTTTTCT	TCCTTCTCTT	CACTTTGGGT	TTAATTTGTT
65551	CATTTTTTT	TCTGGCTTCT	TGAGATAGAA	GCTGAGATCA	TTGATTTGA
65601	ACCTTTCTTC	TTTTCTAAAT	AAGTGCATTT	AAACTTACAC	ATTTCCCTTT
65651	AAGCACTGCC	TTAGCTGTAT	CTCACAAATT	TTGATATTGT	CTTTTCATTG
65701	TCTTTTATTC	AATATATTCT	AATTTTTCTT	GTGATTTCTT	CTTTGGCCCA
65751	TAGGCTGTTT	AGAAATATGT	AGTTAGTTTC	CAAATATTCG	AAGACTTTCA
65801	CAGATACCTT	ACTATTATTG	ATTTCTAATT	TAATTCTGCT	ACADTCCAAG
65851	TATATACATT	ATAAAGTTTC	AGCCTTTTGA	AATGTATTAA	GAATATTACC
65901	AGAGATAAGA	AGATAAGAAT	ATTACCAGCG	ATAAGTAGGG	ATATTTCATA
65951	AATAATAGAC	GAATTGATTC	ATCAAGAATA	TACAACAATC	ATAAATGTGT
66001	ATGTGTCTAA	TAACAGAGTC	TCAAATTATA	TGAAACAAAA	CTGACAGAAC
66051	TAAAGAGAGA	AATGGCCAAT	CCCACAATCT	TTATCTTTAT	CAGGTGATTT
66101	ATCTTGGTGA	ACATTCCTTG	TGCTCTTGAA	AAGAAAGTGT	ATTCTGTAGT
66151	CATTGGGTAT	AAAATTCTAT	ATATGACAAT	GAGGTGATTG	ΑΤΑΑΑΑΤΤΑΤ
66201	TTAGATTGTC	TATATCCTAA	GTTTTGTAGA	ATTATTTCAT	GAATTACTAT
66251	GACAAGGATG	TTAACAACCT	ACAGCTATGA	TTGTGGAATT	GGCTATTTCT
66301	CTCTTTAGTT	CTGTCAGTTT	TGTTCCATGT	AATTTGAAAC	TCTGTTATTA
66351	AACACATACA	TTCATGATTG	TTGTATCTTC	CTGATGAATT	GGTTCCGTTA
66401	TTATTTATGC	AATGTCCCTA	TTTATCTCTG	GTCATATTCT	TTATCTTGAA
66451	GTCTTTTTAA	CTGATATGAA	TGTAGCCACT	TCATCCTTTT	TATGCTTACC
66501	ATTTGCATAG	TTTATATTTT	TCCATTATCT		CTATTTATCC
66551	CTTTATACTT	AAGTCCATGT	CTTGTAGACA	GTATGCAGTT	AATTGTGTCT
66601	TGATTATTTT	TACTCCTTTC	TGACAATTTC	TGCCTTTCCA	TATAATATGC
66651	TTATCAATAC	AGTTGGAGTT	AAATCTACCG	TCTTGTTATT	TGTCACATCT
66701	CCCATCTTTT	GTTGTTGTTC	CTCATTTCCT	TGTTTATTAC	CTTCTTTTCA
66751	GTTATTTTTT	TTTTGTATTC	CATTTTAATT	CCTCAATTGG	CTTTATAGCT
66801	ATATATCTTT	GTATTATTTT	TTATTGTTTG	CTCTAGGGAT	AGCAATATGT
66851	ATACTTACCA	CAGACAATTT	AGAAATCATA	TTGTACCACT	TCACATAAAA
66901	TAGAAGAAGC	TTGCAGCAGT	CTATGTCCCT	TTACACTCCC	ATTCTTTGTG
66951	CTATTGTTTC	CGTATGTATT	ACATCACGTA	CATTGTAAAA	TCCACAATAG
67001	AGTGTTATAA	TCTTTTTCCA	AATCCTTGTG	TGAATTAAAA	ATTTTATGAG
67051	TAGAAAAATA	CATATAACAT	TTTATTCTTA	CCTACATACT	TACCAGTTCT
67101	GCTTTCTTTT	CATTCTTACC	TGTTTCAGTC	TTATCTGTAA	ACCCGTTTTC
67151	ATTTGGTGTC	ATTTCCATTA	GCATTTCAGT	GCAGAACTTC	TAGCAACATA
67201	TTCTCTATTT	CCATGTATCT	TAAAATATCT	TTATTTTGCC	TTCGTTTTTG
67251	TTTATATAAA	TAATTGGACA	TAGAAATCTA	GGTTGGCAGT	TTTCTCTTAT
67301	ACTCTTGGGT	TTCATTGTCT	TCTGATTTCT	GTTGTTTATG	AGGAAAAGTC
67351	ATTGATTATT	TGCTCTTTCT	CTATACACAA	TGTATTATTT	TTCTTTGGCT
67401	GTTTCAAGAT	ATTTTTCTCT	TTATCTGTGG	TTATCAACAC	TTTGATTATG
		TGGTATTATT			TTGGTGTTCC
	TTGAGCTTCT	AACTTCTGTG	AGCTTTTTT	TTCTCAGCGA	<b>ATTTGGAAAA</b>
	ATTTAAGCCA		AATTTTTCTT		
	TTGGAACTCC		AGGTTAGACT		GTCCCATAGA
67651	TCACTAAGAC	TCTGTTCATT	TTTCAATTTT	TTTCTCTATG	TTCTTCAGAT
67701		ATCTTGATCT			TTTATTATGC
	CACCTTCAAT	CTGATATTAA	GGCCATTCAG	ATCTAGAATT	TCTATTAGGT
67801	TATTATTAT	AGTATTAATT	TCTCTGCTAA	GATTTTTTGT	CTGTTCATTC
67851	ATTATGACCA	CAATATTAGG			
	AGTCCTTGTC	AGTTAATTCC	ATCTGAGTCA	TCTTGGGGTT	ATTTTCTATT
67951	GAGTGATCTT	TACCTTATCT	GTCGGTCACA	TTTTTTTCTG	TTTCTTCACA
68001		TTATTTATTG	TTTGCTGTAT	ATTGAAATGA	AATATTATAA
68051	ACAGTATCAA	TTACATTATC	TTCCTTTTAA	GGGTATTGAG	TTTTGTTCTG
68101	GAAGTAGTTA	AATTACTAGT	AGAACTTTTT	GTTCCTGTCA	AACTTGATCT
68121	TATTCTTTGT	TACAGTGAGC	CTATTTTAGT	TTTAAAGTTA	GTCCTAGGGT
00701	ACAACTCTTG	CTCTATTGTA	TGCTCCTTAC	TTCTATCACA	TTTATTTCTA
68231	TTGCCTGAGA	TAGTCAATGA	GTTCTCACCT	GAGCAGGAAC	TGCAACATTT
60301	CARCATGG	TCTTACCTAT	GTATTCATCA	TTCATCTCTC	AGGCCTGTAA
60401	GAAGAGATCT	CTGTTGGGTC	CTGTGGAATC	TTGCTTGCAC	TTGGACAGCT
CO 45 1	CAGCCTTCAG	CCAAAGACTT	GCAGGAAAAC	CCCATAGAAA	CATCTGGGCC
60501	TCTCAATAT	TTGATGTTTA	GGAAGCTAAA	CGTCAAGTAT	AGCCTCCTTT
60551	TOTAGGGACC	CTATCTTGTG	AATTTCACTC	ACCTTAACAA	CTCAGAACTC
60501	TTATCTTCTG	CCTTCTCAGG	GGAGCTAAAC	TGTCACTTTC	TGTGGGCTCC
00001	ATCTTCCTGC	TCCACAATAG	GAAAGTATCT	GCAGAGAAAA	GGCTGGACAA
10000	TCCACECETA	ATTGCTTCAC	GCATTTCCCT	TCTCTCAAAG	ATTGTAAGTT
68751	CCACERROAM	GCTGTTCAAT	ACCTGAAAAT	GATTTCTACA	AATTGTTTTT
69001	CCAGTTTTAT	GATTGTTTTC	AATGGGAGAT	CATTTCTAGT	ACCAGTTCCT
2000T	ACTOCCOCAT	CAGAGGTACA	AGTTCAACTT	GGATCATTTT	AAAAATACAA
68001	TCTTACARTA	GTCACTTCCT	GUCCUAAACC	CCTTGGTAGC	TTTCCATTGC
68951	TOLINGAATA	ACTTTGTGAT	CTACAACATC	TTCTTCAAGG	CCCCGCATGA
69001	ATCCACCOMM	GGCTATTTCT	TOTAGTTTCTT	ATTGCACCAC	CITGTCCCTC
69051	TTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTT	TTTTTAGTCT	TCTCTCTTTC	TTTGAACTTC	TACCACCAGG
69101	TCTTTACACA	CGTTCTTCTT	TCCCCATTAA	CAATGATCCA	CCATTCTCTT
69151	ATGGCCAMMC	CTGTTACTCA	CACTCATAAC	TGAAACATCA	TTTCCTAAGG
69201	CTTCTTTTT	CTGGTTCAGT	CAGTCTATAT	CACCCCC	ATCACATACT
69251	ATGTGTTTTMC	CCTATATTTT TATTTTATGT	CTCTTCAAAG	CACTTATTTA	AGTTGTAATT
JJ2J1	GIGITGIT				CACAGTCCAG
			T ~ T TT 7		4.0

FIGURE 3, page 18 of 57

69301	GAGAACAGAA	ATCCTGCCTC	TTTTATTTAT	ACCACATCCA	CAGTATTATT
69351	AGTGCCTGTC	ACCTAGTAGG	TATGCAGTAT	GTACCTATTG	AATAAATGAA
69401	TTGACTTCTG		CGTCTACTCA		
69451	ATAATACCTT				GAGGATTTTC
69501	TGCATAGCTC			TTTAAGGAAG	ATTGTGAATA
69551	TTATCAGATA	AAGATTGTGA	GACACAGAAA	AGCCAGATGA	TTTGGCAATG
69601	CTCATAGTAC	CAGAGGCAGA	AATACAGCTA	GAACAGTCTC	CTGGCCTCTA
69651	ATCAGGAGTT	CTTTCCAGAA	CACTGCTTCA	TCTTCCATTC	TCTTGGGTTC
69701	TTTCTATCCT	TACTTTATAG	GGCAAAATGT	GTGCAAAGTA	TARTCCCTCT
69751	TTTGCAATGT	GTTTTTAGTT			GGCTTTTTAT
69801					AGGGCTTATA
69851					CCATTTCCAT
69901	TCTTGGCCTT	GTGAAGCTCT			
69951			ATGCATCTTA		
70001		AATAAAGACT	ACGTCAAAAA	TCACCCATCA	ATCGACAACC
70051					AATGTGCTGA
70101	GCTCTGCATT		CTCCTCTGTG	ATCAGGGTGG	ACATTCCCAG
70151		CCGAGGCTGG	AAACACCATC	TGAATGTCTG	ACCACACAAA
70201		TGATCCAGGT			ACCACCTTCT
70251	AAGCAACACT		AGCCCAGTTA	ΤΥΤΑΤΤΟΚΑΟ	GGGATGATTG
70301		TAGTGTTTCT			TATTTATCCA
70351			AGGAGAACAA	GGTAGGCCAA	ACTECCTTTE
70401	TACTATTAAA	GGCTGCTTGA	TTTCTAAGTA	CATCTTCTT	GCCACCTTTC
70451	TGCCATTCCA	CATTCTAGAA	GCCATGGGTA	AGTCAGCACA	GCCACCITIC
70501	CATGATAACA	TTGGTTTTAG	GAGGTCTCGT	GCATAATGGA	CCAGACTTAG
70551	AGCACAATGC	TGTAAGGTAG	TGATTTAGGT		
70601	AGGAGTTTAT	TATCAGATGC		ACTTGTGGCC	
70651			AGCCTTAGAA	CTCTGGGAAT	TCTGAATATA
70701	ATTCCTGAAT	CAATCGTAAG		TGATGCTTAG	TGCAAACCAA
70751	GAGGCAGAAT		AGTGTATCCT		
70801	TTTTCCTGCC		CTTACTTTTC	CATCCTTCCT	CATCCTACTA
70851	CTAACTACAT		TTTACGTGGT		CTAAGCTGTT
70901	AGCTTCATTC		AGGCACTCTT		TACAATTGGG
70951	AAAACTGAGA		TAAAGTAAAT		CATTATGCTA
71001	GTCCATGAAG		TTGCAACTAA		
71051		ACCAGAGGTT	AGCAAACTAC	TTCTGTAAAG	GCDACACACT
71101	AGTTATCTTA	ATCTTTGTGG	GCAACATAGG	GTCTCTGTAA	CCTATTCTTC
71151	TTTCTGTCAC	AATCTTCTGG	AATGTAAAAA	ACATTTAAAA	TTTACAAACC
71201	TTACAAGAAC	AGCTCATGGG	CTAAATCGGA	CCTGGATTTA	GTCTGTGAAT
71251	CATAGTTTGC	TGACCCCGCT	TTTTAACCAG	TATGTACCCT	CCTTCTCGGG
71301	ATGTGAAAAA	TTAGTGCAAT		AATAGCAAGA	
71351	GGCCTGGAAG		ATTACATCAG		
71401			GAACCACCTG		
71451	CTTTCCCTCT	GAGATTCTTT	CAGGAACCTC		CAGCCCGGAG
71501	AACCGTGGGC		TGCCTCCTCA		
71551	GATAGAAATC	ATCACATCTG		CTCAACCTTT	CTCTTCTGCA
71601	CTTTCTTGGA		GCACTACCAG		
71651	TTTCATTTTT	GTGTTTTCAT	TTAATTATTT		AAGTGTTTGA
71701	CTGTTTAAGG		GTAAATATTT		TGGCAGAAGC
71751	TGTGGTTTCC	TTTGATGAGC		TGGCTTTTAA	
71801	ACCAGGACAG		CCAGTGGGTG	CAGTCCCCAG	CACTCCCCTC
71851		AGAAAGCGCT	GCTGGGCCAA	CAGCCTTCCT	CABCTTCCC
71901	GCTGCCCCCA	TCTAACCAAC	ACCTCAGTCT	CTTCTCCACC	TECTTCCCTE
71951	CCCTCTTCCT	TTCCCTCGCA	GACACTTTCT	TCTGCCTGGC	AAAAGGAATC
72001	TTGTTTCCAT	GGAAGCCTCA	TTAAATCTGC	ATCTTGCTCA	GTTTGGGTTT
72051	GATCACGGCT	GCCAGAAGTA	TTTTTAGCCC	ATGCAGTTGC	GTAATGAGAT
72101	AGAGATTGGG	GAAAGGGGGA	GGTGACTGTA	TAGGCAGAGG	GTTTTTTTAA
72151	AAAAAGTGA	GAAAGAGAAG	GAAAACCTCT	AAAGAAAAGA	GTTTTATGGA
72201	ATTGGAAGAA	GGATGGAGCA	CCTCTTTTGG	GAGCATGAGG	CTGGTGTTCT
72251	CTGGTTAGCT	CTTCCCACTG	GAAGCCCATG	GACACTTGCC	ATAATACCTC
72301	TCCTGGTCAC	ATGTCAGGGG	AACCTCTGAT	CTCCCTTTCC	ATGAGCTTAG
72351	TTGGCCCAGC	CAGGGTGACA	CTTATGCTAG	GGAGTGTGAT	TCATCTTCCT
72401	GCTTACAGAT	TTCCCCTCCC	ACAGACCTGA	TEGEGECAGEC	AGGATAGTCG
72451	CAGAGAAGAA	GACAGAGCAA	TAGCAGGAAA	GAGAGGACAA	CACTAACACA
72501	TTGGAGGTTT	ATGTTCAAAG	ACGGGATCTA	GGGGGTCAGA	GAAAGCACAC
72551	CTACCATGTA	ATTGGTGCTG	GAATCTGATG	CCAAGTGCAC	CCTTGCCTTC
72601	TGAGGTTCTG	AGAACTCTTG	CTTGTGCTTT	TCAGCCAGAC	TATECCCCTCA
72651	CCTGCCCCTG	TACTTTAAAG	AGCTCTTTAG	GCTGGAGTGG	TTCTTTCCAT
72701	TGGATTGTTG	GAGTGTGTGT	GCATGTTGTT	CTCTGGGGGG	TIGITIOCHI
72751	AAAGAGATTA	AAAAAAAACC	ACATGCAGCT	GTCACACCTA	ATTACHAGEC ATCTTTATATT
72801	AACTTTTACT	ATGCCACATG	GTGTTTTAAG	CATTCTATAT	GTGTTANTIG
72851	ATTTTCCCTA	ATTCTATGGA	CTAGACACTT	DADCACTOTA	CATTCTACAA
72901	ACAAGGAAAC	TGAGGCACAG	AGAGGTTGGG	ADDUCTURE TO THE PROPERTY OF T	CALIGIACAA
72951	AGCTAATTAA	TAGTGGAGCC	AGGTTTTGTA	CCCDCDCDAC	CTCATTTCAC
73001	AATCTGCAGT	CCTAGATTAG	TAACGTGTTG	TTGGCCTCTC	DCDCDTTTTA
73051	AATGACATTC	TGTACACAGA	ACCATTTATA	GTAACTTTCT	ATTCTTTIA
73101	TGAAAGCAGT	CTGCAGATGT	GCTGCTGGGA	TTTCITIOI	CTTCNANCAC
				Chilchi	CIICAAAAA

FIGURE 3, page 19 of 57

73151	GTGTTTTTT	TTTTTTTAA	AGGAAAATGC	TTTTCTGAGG	GTGGTATCTA
73201	AATTCATAAA	AATCTTTACG	ATCAAGATTT	TCACAAATTT	CATTCTGACT
73251	CTGTTGCATT	GCCCTTCTTC	CCATATTCCC	AGTTAGTTTG	TATTGATTGC
73301	TGCATCTCCC	TTGAGCCCAT	GGTCCCCCAC	A)C)	GCAGAACTGT
73351		CACACTCTCA	CCCACCACCA	COCMOMONIA	CGGCCAGCCC
73401		ACCTCCTTCA	TCACCA CCAC	GCCTCTCTAG	CGGCCAGCCC
73451		AGCICCITCC	TCAGGACGTT	TAATTTCCCA	CATTTCTATG
	CAGTTACCTC	ACAGAAGGAT	GGCTACGAGG	GCCTCACTTG	GCTTGGCAAG
73501	TTGGTCCCCT	TTTTACTCAC	AAGACTCTGT	TTATCTCTTT	CTTTATCTTT
73551	GTTTATCTCT	TTGTTGACCT	GCCCCTCTTC	AAGGCCTCAG	TTTTCTCTGA
73601	AGTTTACAGC	TTCCCTCCTC	ATCCCGCAAA	AGACCAAAGT	GGAAAAGATG
73651	AAACCAGAAT	CCACTGCAAG	CCCCACCTGC	CACACCCTCT	CCTTTTTGTTG
73701	CATTCTCTGT	TGTGTTTAGG	ACTTGAGAAT	CANCACCICI	ATGAATTGAG
73751		TTATTCTTTA	CAAMAMCCOM	CMAGAGGGAC	ATGAATTGAG
73801	ACCCCCATT	CATACAMCAC	CAMIAICCCI	GTGAGCTGAG	TACTGTAAAT
73851	MCCCCCAIII	GATACATGAG	TAAACTGAGG	TGTGGAGTGA	TAGAGGAATT
	TGCTCAAGGT	CACATAACTA	GTAAGTGGGT	GGAGCTGTGA	TGTGAAACTG
73901	GGCAGTCTGA	TTCTGGGACC	TGTGCTCTTA	ATCACCAATC	TATATTGCCT
73951	CCTACTTGAA	AACATCCAGG	GAAAATGTTG	AGATAGATCA	GCTGAAATCT
74001	TCTTGCACAG	TAAAGCAGGG	GCCACCTGTC	CTGGAGTTAC	ATTCATCTTC
74051	TTCATTGTCA	ACGATTTGTG	TTCAGTGACA	CCCTCTTCAG	CCCAAGAACT
74101	TACCTGGGTG	CTGTGACAAT	TGGACATGAC	TACCAACAAC	CACTCACATTC
74151	GTAGCCCATC	CAAACACAGG	GTAGGAAGTC	CAMCCAMCAMC	CAGIGACAII
74201	GGTTATTAGA	ACCACCAACC	CACMANACCO	CAIGCIIGIC	ACTOTOTTT
74251	CTTCAATAGA	AGCAGGAACC	CAGTAAAGGC	ACCTTTTATA	TATCTATAAA
	GIIGAAIAIA	TAAGATATAT	GGGGGCCAGG	CACAGTGGCT	CACACCTGTA
74301	ATCCGAACAT	TTTGGGAGCC	CAAAGCAGGT	GGATCACCTG	AGGTCAGGAG
74351	TTCAAGACCA	GCCTGACCAA	CATGGTGAAA	CCCCATCTTT	ACTAAAAATA
74401	CAAAAATTAG	CTGGGCGTGG	TGGCACACAC	CTGTAGTCCC	AGCTACTTGG
74451	GAGGCTGAGG	CAGGATACTT	GCTTGAACCC	GGGAGGTGGA	GGTTGCAGTG
74501	AGCAGAGATT	GCGCCACTGC	ACTCCAGCCT	GGGTGACAGA	CCCACATTCC
74551	ACCTCAACGA	АААААААА	CDDCDTDTDT	CCCMPMCMCM	BCARCHCACA
	GAAGGGCAAA	CAGGCCTTAA	CACCACCACA	BREARISTO	AGAACTCACA
74651	CCACTACCTT	CCTCTCTTTTT	CAGGIGCIGA	AAACAGGAAC	TGGGAAGTTG
74701	CCAGTACCTT	CCTGTCTTTT	CCCCTGGAAC	CAAACGGTTT	CTTACTTGCT
74701	TCTCTCTGCA	CCTCTGTCTC	ATTTCCCTCT	CTCTTCAGAT	GATTTTTCAT
74751	TGTTGCATCA	CACACATAGA	AAAATCAGGA	TCCACCCTCC	CAAGTTTACA
74801	TATCGTTGTT	TCAGGCAGCC	ATAGTATCCT	TAAAACTCCA	CATTCCAGGG
74851	AGAAAGCTTG	GGTCAAGGAT	TCAGCCAAAG	GGCAGCGAAA	TEGRETARA
74901	ATGCAACTGC	CAGGTCTATG	GGCAGCAAGG	AGGCCGGGAA	GGAAGCCGCT
74951	GTTGTGGTCC	AAGTGACAAT	TCDACAGCTC	AAACCATAAC	TARCOCCCC
75001	GCTTTTCACA	GATGGAGAAA	CTCACCCACA	CAACCAACCO	CCCCCCCCCC
75051	CAGGTCTCTG	CCCTTTCTCTCT	CAAMCCOACA	MCACHACCI	GGCTGGGGTC
75101	TTTCTACAGG	AAATCTCCTT	CAAIGCIAGG	TCACTGGATG	TGGCGTCTGA
75161	TTTCTACAGG	CORCOGGII	TCTCTACTTT	GTCCCAGAGC	CCACTCAGAG
75251	CACTGGCTGG	CCAGGGGGTC	CTAGGGCCCT	CTTAGGATAG	TCTCAGGCCA
75201	ACAGCCCCAG	GACAGAAGCA	ACCAAAGTGA	AGTTATGAAA	GAAAGCTCTT
75251	TGCTGATCTG	TCAATGGCAC	CCTTGTAGAG	CCAATACTTA	GAACACCTGG
/5301	ATTTGAATAC	TCATCTCCAA	AACCTGTGTT	CTTTCTACCA	CGTGACAAGC
75351	CCTTGTAAAC	CTCACAACGT	CTCTATGAGG	TGAGCGCTTG	CAGATCCACA
75401	CTTTAGATAA	GCAAATGGAG	GCTCAGAGGG	TAAGCAGCTA	GTTCAAGGTT
75451	ATGCACCTGA	GCCAGGATGT	GCACACACCT	CTCTCTCTCTA	TTCCTAGGII
75501	CCTGTGCTTT	ACCCACTTTC	CAATACTCCT	COMMENCE	TICCIAAGGG
	ATCTCTCACA	TCCCAACCAM	CAMIACIGCT	GCTGTCTGCT	TCATTTCCTC
75601	ATCTGTCAGA	TOGGAACGAT	AATACTCAAC	TCACATGGAT	ACTGTATGAG
75651	CORCARCOS	TAAAAGAAGA	GAAAGTGCTT	TGAAAACATA	AGCAGCCCTG
	GCAGATGGGA	ATTATTTTTG	CTGCTGACAC	ACATCCTCAG	CCTTGAGGGC
75701	TCTGCTGAGC	CATACCCAGC	TCAGAGCTCT	GGAGGCACCT	CCTCCCCATC
75751	AACAGCAGGG	GGGACATTCT	GTCTTCATCC	TGAGCAGGCT	GACAAACTGA
75801	ACCCCACTCC	TCCCTCAATG	TCCCCATGCT	GGGAAGGAGT	ATAGCTCATG
1282T	CTGTGTTCTG	TCTTGTTGCT	GAGAGAATGC	AGAACCCAGA	ATTTGGGTCT
75901	CAGCAGGTTG	GGGAGAAAAG	GAAATGTATT	TCTTCCCCCA	AGATTTCTTT
75951	TTGAAATATT	TTCATTTGTG	GAATCAGATT	GTGCATGCAA	GTTTCTTCCA
76001	GAAATGTAAG	ACGTCGTAAT	GATGGGAACT	CTTCCTTTT	TAATTCAACC
76053	ATGGGAAAGG	AAACTGATAT	TTATECACCA	CCACAACAA	DCCACCCACC
76101	TACCCARCCA	TO TO TO THE	TINIGGAGCA	CCIGIICIAI	ACCAGGCAGC
76161	TACCCAACCA	CMCCATTG	TTGCAATGTT	ATGCAAGCTT	TATTATCCAC
70131	ATTTCACAGT	CTGAGTCTGA	CTCAGCAATG.	TTGTGTTCTA	TGTGCTAGTT
76201	CCCACAGGTA	GGTGGCTGCA	GCGCTGGGAT	TTGAACCCAT	CTCCAAAGCC
76251	TCCATCTTTC	TACCACTGCC	TCCCATTGGT	GGGGAGGCCA	TGGACTGGCT
76301	GTCAGAGATG	TCCTTTCCAG	TCTAGCAGAC	TAGGAAGCTG	CTGGAAGCTA
76351	CTTATGCAAA	GGTCAGCAAG	GAAGGAAACA	GAGTCAGAAC	TAGATGGGGC
76401	TCCCCTGGCC	ACTTTTCCAT	GCTGGCCCAC	ATGTCCGGCT	AGCAGTCAAC
76451	ATTGGGTCTT	ATGCAGAGCC	ACCTGTGTTC	PATCENTACA	TCCTCCACAC
76501	TGCACAAACT	AGTGGGACCC	TCTCTCTCTC	CACCOMOMOCA	COMMONWE
76551	GGTTCAGCCC	ABCTCATCAC	CENCCCCNCC	MACCA CA TOTAL	GGITCATTGA
76601	CCADATCCCC	CACCUATGAG	CIAGGGCAGG	ACCAGAGGG	TGTGTTCCAC
76661	CCAAATGGGG	AGGGCA AGGCA	GGGGACACAG	GCTCCATTTT	CATGACCAAA
16021	GACTGAGCAG	AGAGGCTCTC	TGAGCAGTGG	CAGAATGGGA	AGTGTCAAGA
10101	AGCTTTGTTT	GACAATTGAG	TCAAGAGGAC	AGAAAAGACA	GAAAGCAGAC
10/51	ATCAGAGTTG	GGAAGGCTCA	CCCCAGCTCC	TTGACAAAGG	TGCATGAGGC
10801	CAGTTCTTGA	AGCAGTGACC	CTGCCTTATG	TCATGTGTTT	ATCAAAGCCC
76851	GCCCATCAGC	CCTGAAGTGG	CCTCTGTGTT	TAGAAGAGGG	CCTGACATCA
10901	TTUTCTGAGA	AAGGATTTGA	CAACAACAAA	GTGTTGCCGT	ATGTGTTGTC
76951	TCATCCCCTC	AATAGTCCTC	TGAGGTATGT	GAGACAGGTG	TTACTCTCTC
					- ARCICICITY
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FIGURE 3, page 20 of 57

77001	CACTTGGCAA	ATAGGGAAAA	GAGGGCCCAG	AGAAGTGAAG	CTGCTTTCCC
77051	AGGACCACAC		CAGTGTCCAT		
77101	CACCAAATAC		ACGCAAACAC		GGACAACCAA
77151	GTCATCTAAT		GCTATGGTCT		GTCTTTCAGG
77201	GCTATACCCT	AGGAGAGCTA	ATCATTCTTG	GTTAGATAAG	AAATAGCCAA
77251		GCATGGTAGG		CCAGAATAAA	
77301	AAGAGATGCT	CAGAATGTGT		CTTCACTATA	
77351	GTATAAGCCT	TGTCCATCTG	TCACATTATG		GCTCCCACCT
77401	CAATTCCTGA	TTCCACATTA	CAACAAATAC		TTTGAACTAA
77451	CAATGCCAAT		CCCATATTAA		CTGAGTCAGC
77501	TACTGGAGGT		ATAAGATGGT		TTAGAGGATT
77551	CTTTGGTTGC		CACCCAGCTT		TGGAGAAATT
77601	GGGATTTTTT		AAGCAAAGCA		
	ATGATGAAAA				CTTTTTTCTT
77701	TTAGGCCATC		CAAACTGGTT		GGCTGGAGAC
	GTTACTACCA				
77801			AGGCCATTCA		
77851	ATGTAAAAAG	ATAGCCACTC		AATGTGGTTA	
77901			GGGGTGTGGG		
77951	TGATAATACC		GGAGCTTTCT		
78001	ATGGACAAGT	AATCAACAGC		AGCTGCCAGC	
78051	AAGAGCCCTG				ATGTGGGTCT
	GGATTGAAGT		ATCAGGAAAG		
78151	ACACTTGAGC		AGAATGACTG		
78201	GGAGGAAGTG	CACCCGAGAC		ACATAAGCAA	
78251	GAATATGAGG	ATCGGGGAAG			
	GGGACACGAA				AGAAAAATCA
78351	AAGTCCTTTG	AAAATCATGT			
					CTGTGAGGGT
78451			GAGAAATGTT		
78501	ATAACACAAT				GAAAATCTCA
78551		GTAACTAATG		TACATTTAAA	
78601	ATGCAATTGT	GAGGATGATG	GTGAGATGAG	CAAAATAATC	CAGTTTGTAA
78651	TTGTAGTTAT	CAGGCTGGCA	TATCCTGCAG	GTCACACTTC	TAAACATGAC
78701	TTCGAAAAAT	CAAAGATCAG	CTAAGTTTGA	AGTAAGTATT	GAAAGAGGGA
78751	GATTATGTTG		AAATAGAACG		
78801	GATCAAAAGC	ACCAAGCTTC			GGGTGCGTGG
78851	CTCCGACACC	AGATATCTGC	AAAGCAATAT	GAAATGAGAT	CAATAGTAGA
78901	CATTGAAAGA	TTGAAACTGA			
78951	AATGAAATGA	GACCTAATAA			
	ATTAAAGAAA				
79051			CACAGGAGCT		CCATTCTCCC
79101	TCCTGCTAAA	AGCCGAGTTT	GTTTTAGCTG		
79151		AAAAAGGAGC		TGGTTTCCAT	TATAAAATCA
79201	GAGCTCTGCT	GCCATAAAAT	TAAATCCCAT	AATAAAATGA	GTAGAAAACG
79251	TGATGTCCTG	CAGAAAGGAA	GATGGCAGCC	CACTCAGTGC	CATGCTGGGC
79301	TTGACTATAT	ACAAGCCGTG	CATCTCCTGC	TGCGAGTTGT	AGCTGCTGCC
79351	CAGCAGTGCA	CATTATCGTT	GCAGCTGTTT	TCCTCACATT	CTGAGGTTTA
79401	TGAAATCCCT	CATCCATCAA	TAATTGATCT	TTAGCTCTTA	GTCCAGGGGT
79451	TGTCAACTGG	CACTCCATGG	ACCTTTAGAG	GATTGATGGC	TAGGTTTTCA
79501	AAGATCTTTG	AACCCCCTGA	AATTATATAC	AAAATACTGT	GTGTGAGTAT
79551	GTGCATTTTT	CTGGTAAGAA	GCACCTGAAT	TATCGAAGCA	GTTTGTGATC
79601	CCCCAAAAAG	CTAAGAACTA	CTTCCTAGAG	CAAAGGGAGA	TTTTGCTACA
79651	CTTAGAGATT	TACACATTTG	ACCAGGGCAG	CTCACACAAG	TGGGATGCGG
	TTTCACATTT				
79751	AATATTGTAA	TACTTCTATA	TGAATCAGGA	ATTCACTATA	TTTAACTTAT
	TTGGAATAAG				
	TTCTTCCTGT				
	GTTTATATTT				
	GTATACCACG				
	ATGTGACCCC				
	CACTGAGCTT				
80101	GTGATAGTTT	AAGTACTGAT	AATTATTCAC	TTGGAAGGGA	<b>AGAGAATAAA</b>
80151	ATTCAGAACA	CAGTATTCCT	TAATGGGAAA	TCAACTTAGA	<b>GGAGGTAGGA</b>
	GGGAGATCAA				
80251	AGCCAATGTG	TTGATCAAAG	AAATTATCTT	TCGGGGAAAA	CAGTAGAAGG
	CAATTGAAAA				
80351	AATGGCTTGA	TTGTGTGATG	AGGTAATTAA	TGGCTGCAGT	TAGCAAAATA
80401	TGTTCAAAAA	AAAGACAGAA	AGGGTAGTTA	CAGGAGAAAA	<b>ACATCCCCGC</b>
80451	AGATCTTCAA	AATCAGAAAC	AATGAAAATA	ATTATTTCAA	AAATTAAGAA
80501	AAAAACTCTC	TAATTTATAC	CTGAATTACC	TGGATAATTG	GTAAAATTTC
80551	CTGCATATAC	AAATCTTGGT	CCTCTGCTCC	TCTCTCTATA	AATAAATAGA
80601	AATGTATGAA	TCAATAGTCA	GCCAATGTGT	TGATCAAAGA	AATTATCTTT
80651	TGGGGGAAAA	TTGGTAGAAG	CCAATTAAAA	AACAAGCATC	ATATTGCATG
80701	AAAACAGCAA	ACGGAAGTCA	CAATGGCTCG	ACGGTGTAAT	GAAGCCACAC
80751	AATATGTATT	AAACACATCA	TCTACACAGA	TGGATTCAAA	GATACCTTCT
80801	TTGTGTCTAA	GTCCCAAATC	TGTGTTTCCT	GGCTCTGTTC	CCTCATATCT

FIGURE 3, page 21 of 57

80851	AGTCATTCTC	CAAGTCAGCA	TGCCCAACTT	GAAAGTGTCA	TTTTCAAAAC
80901	CTGCTTCTTC	TCTTCTGGAA	GTTCTTCCTC	TGCCCATTGC	TCCACAATCC
80951	CCACCTCTTT	CACCCAGTAG	CAAACCTTAA	ATTTATCTTT	TACTTTGTCT
81001	TACTTCCCCT	TCTTATATTC	AAAATGTTTC	TCACTTGCAT	
81051	TCATTTCATA	AGCATTTATG	AGCTCCTGTT	ATGGTTTGGA	AACTCTTCTT
81101	CATGCTGGAG	GTGGTCTTAT		TTTCAATTGA	
81151	TGTTAAGTGC				
81201				GCTGTGGGAG	GCTCCCCAAA
	TCAGICIANG	GAAGTTGGGA	AAAGCATCTC	AGAGAAGATG	GTGTCTGAGA
81251	TGGGGAGGAT	GTGTGGAACT		GAGAACAAGT	AACAACATTC
81301	TAGAAAAAGG	CCTCTTTCAG	CATGCTAAGA	AGTTTGGAGG	ACAGAGGAGT
81351	TACCATTCAA	AATTTGGAGG	GAAGGAAGAG	CATACTGAGG	TTTGCCACTT
81401	GAACAGATAA	TTTCAGCTGT	GTTGGGTGAG	TGAAGTTGAG	TGGGTACAAA
81451	TCAGGTCAGG	AATATAAGTT	AGGAGACTGT	TACTAGAATC	CAGGCCAGAG
81501	GTGATGGTGG	CCAATATATG	AGAGTTTTAG	CAGGGAATGA	DADADACAAA
81551	ATGTGTTCAT	GAGGTAGAAG		CAACAGGATC	
81601	TTGGAAATGG			CAGAATGCAG	TGGTTCCTGA
81651		CATCCACTCA	CAGAGGAAGC	CAGAATGCAG	
81701	CCTADACAAA	CATCCACTGA			AAGAAGCTTT
		ATAAGAAGTT		ATGTTTGAAT	TTGATTTCTC
81751	TGATGAGGAG			GCTCAGGGTG	TAGACTTGAG
81801	AGTGGATGGG		GTTGAGGCTA	TTAAAAGGGA	AAAGGTCAAG
81851	GAACTGAGGG	CCAAGGATTT	ATAATAAGTT	<b>ATCTTGGGCC</b>	ACTAAAGCCA
81901	CGCAGGATGC	TGGCAGGAAA	CCTATGAGCC	AGGTCTTCAA	TGTTGAGTCC
81951	AGTGACTCAG	GTGTCAGAAG	CAGCAGGAGA	AGCATTGATA	GCCTGATGGG
82001	GAAGGAGCCG	TTACCTGAGA	GTAGCAGAGA	GAGTTATCCT	AGCTGACACA
82051	GCTCTCAGGG	ATTTGCTTCT		TTAGGAAAGA	
82101		ACTGGTGGGC	ACTCCCTTCC	CCACARARGA	MAGAGCAGTA
	AATTCTATTC	TCAACCCACO	ACTOGCTICC	CCAGAAAACC	
82201		TCAAGGGACT	CCTATTTAGA	TAAGGGGCTT	TGTTAGTTCT
	TOTAL	CCAAACAGAT	GTATATCTCA		
82251		CCACATGGGC	CTACCCACTG	TCTGCTAAAT	GCACTTCATA
82301	TTTTCTTGTT			ATCTTCTTTT	CCTAATCTCT
82351	GCCCCTCACT	TACCTGAATC	TTTTGTATTC	TCAATGACCT	GCTCCATCCC
	AGCCCTTTCA	AGAACCTTTA	ATACCTACCA	AGTGAATACT	CTCTCCATTG
82451	ATTACACACT	TCCTGTAGCA	CCTGTTCTAT	AATTATGAAA	TATTACCTAT
82501	TGTACACATA	TATTTCAATC			AATTTATGCC
82551	TTGTCAATTT	GTAGCACATT		TGTAGATGCA	
82601	TTAGAGAACT	TGTTAGTTAA		AACATGGGCT	GCAAAGTTCT
82651	GGTCCATGCA		AAAATAGAAA	TITOLITOGGCI	
82701	TTAAATTCCA	TGAAGCAGAA	λελπλησολελ	CAMCCACCAC	TGCATGGAGC
82751	CCTGTACAGC	TOTALOCAGAA	ACMINICAGE	AMMOONGCIG	AATTTGTTTG
82801		TCTTACAGCA	CCTCARAGE	ATTIGITIGA	TTTACCTAAG
	AGCTAAAATT	GIMANIGGCA	GCTCAAATGA		
82851	ATGAGTTTGA			GCTTAAGACA	TGAAAAAGGG
82901	GGAGATTATA		CTTTTTTATG	GCAGAGCATT	AAGGAAAAAA
82951	AAGTGCAGAT	AAATGAGATC	AAATGGCAAG	TGTCTGAACC	TGCTGGACAC
83001	AAGTCCCGGT		AGACAGTGTT	TATATGACTT	CTGGGCCATC
83051	AATAGATAGA	TAAGGTACAT	CAGCGGCCAA	TGTTCCAGGA	AGTTTGAGAA
83101	GATAAATGGA	AGTTGCACAG	CAGCCTAAAA	GCTTCCTTAG	GAGGGCTGTG
83151	CTCCTCCAGA	GCGCCATCTG			TTCTTCACAT
83201	TAAATGCTTT	TCCTTTTCTC			AAAGATATGC
83251	TAGCCTGGAC	TTTGACAAGG	ACATCTCCAC	ATARCARACA	TTCTGAATTA
83301	TTTTTCCCTT		TAGCAATTTT		
83351		AGAATATTTC			TAGATGGCTA
83401	GTGGTACCTA		TITICITGGA	AAATCATAAG	GCTTTGGATA
		TAGAAGCTGA	CATCAGCAGC	AGCCTGCCTC	CAGTCGATCA
03431	GGGCCTTTGG	AACTICACGG	GGCTCCTCTA	CTGACAGCCC	CATCGGTTTC
03301	CCTCCAGCAC	ACGTAACTCA	GCATTGACTC	TGGGTAGTAG	AGGGTGGTTT
93331	ATGGAATCTG	ATTCATCTCA	GAAAGAGGTG	GATGCAAACA	CATTCCCAGA
83601	GCAGAAGGCT	TGGCATGTCT	GGTCTTAGGC	AGAGGGAACT	GGAGATACTT
83651	GTCCTATTGT	TCTTGAGATT	CCAGCAAAAA	TAGCCCATTA	CAGAGGAAGA
83701	AGATATCAGG	TCAAATGAAG	GCTTTGGTGC	TACAACATTG	TCTTAGAAAA
83751	AAAAAGAAAG	AAATTGGCCA	AGTGCAGTGG	CTCAGCACTT	TEGGAGGCTG
83801	AGGGGGCAG	ACCACTTGAG	ATCAGGAGTT	CGAGACCAGC	CTGGCCAACA
83851	TGGCGAAACT	CCGTCTCTAC	CAAAAAGTAT	TABABABTAC	CCCACTCTCC
83901	TGGCGGGCTC	CTGTAATCCC	ACCTACTCCC	CACCCTCACC	CCCCACAAMC
83951	ACTTGAACCT	GGGAGGCGGA	GGTTGCAGTC	ACCCA ACAMO	CECCCAMMCC
84001	ACTCCAGCCT	GGGCDACACA	CECACACECC	MUCCHAGAIC	GIGCCATTGC
84051	CDDDDDDDCDD	ADACAAAAAA	GIGAGACICC	ATCTCAAAAA	AAAAAAAAA
84101	GAAAAAAGAA	AAAAAAAAAAAAAAAAAAAAAAAAAAAAAAAAAAAAAA	GAAAAGAAAG	AAATTAAATT	AAAAAAATTG
04101	TTTTTTAAAC	MAAGGAAGGC	TTTGGGCTTG	GAGTCCAACT	AAGCTAGGCT
04101	GGAATCCCGG	TTTCATCTCG	CTTCTCTGTG	CAACTTTGGA	TTTTACTGAA
84201	TCTCTCTTAT	TCTCAATTCC	CTCCTCTGTA	AAATGAAGAT	AATGCTAGTA
84251	CCTGTCTCAT	CAAGTTGAAG	GAGACTTAAA	TGAGATGTGT	TGAAAGCATT
84301	TAGCATAGTA	TGTGGCACAT	AAAGAACACT	CAATAAATGC	TGGCTATAAA
84351	GAAGCCAGAG	AGAGACTCGG	AGGTGATGAG	AGAGGCCACA	ATTCCCTCCA
84401	TTTCATTGAA	AAGCAATTTT	TATTATCTCA	TTTGAAAGGC	AGTATAGTAT
84451	AGTGGTTAAG	GACATGCACT	ATGGAGCTAC	ACCTCCTCAC	ተፈር ፓር ተስታውር ው
84501	GTCTCTATCA	TTTATTACCT	GTGACTTAAC	Chacacacac	CACTORION
84551	ATCATTTTTG	AGAGAGGAGT	שמאליים	CCWVCWCWCC	TO CONCERN CON
84601	AGATTTGATG	AGTTAATACA	TATANACCAC	COLUCIOIO	CCCMCCT CCC
84651	TATTAAATGA	CATCTARCE	TTACCECCEC	ACATAGTAGT	GCCTGGAGCA
J.JJI		CHIGIAAGTA	LIAGCTGTTA	AAATTATTII	CAACATGTGG
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FIGURE 3, page 22 of 57

	•				
	CATAGGACAT				
84751	AAATCAGAGA			TACATGTCTG	
84801	TGGACTGCGA			CATCTCTGCA	
84851	ATCGTATTAT			TGCCAAATGG	
	GTTGTAGGGA			GCAGGCGGG	
84951	GGTTAAAGGT			TCACATTAAG	
85001	ATTGAACTGC			AATCAGGTCA	
85051		TACCAGTACA ATTGTATGTG		AATGCATAGA	
85101 85151	ACTATTCCTA			AGTAAATAAT TGTTTTGTTT	
85201		GTGCTAATCT		CCAACACCAG	
	CACCACAAAAAAAAAAAAAAAAAAAAAAAAAAAAAAAAA			ATGAATTTGG	
85301	TTCTGGGTCA			ATGTCATGTG	
85351		AATGTCTTAC			
85401	CCTATCACAT	ACTTTGCCCA			
85451		GAGGTCCAGT		TCCATCCTTC	
	CAGTTTGTCT	•		CATCCACCTT	
85551	TTACTCATTC	TTCCATCACA		CTCCCATGTC	
85601	CAAGTACCAT	TTGGGAAACA	GAATTCTAGG	AATCTGGAGA	CCTAGAGCTC
85651	TTCAGACCCT	GAAATCCAGT	TTTCTGAGCT	GAGACAGTTT	CTTAATTTCT
85701	CACTCCAACT	CCGTTTCTCC	TCTTTCTCAA	TGGATATTTT	CCAAGTCTCC
85751	ATTAGGCATA	TAGCAATTCC	AGAAAACATT	CAATTTTCCC	TTCTCTTAAT
85801	GCCATGCTCC	AAAACACCAC	ATTCCCTCTA	GACATTGAGC	ATTGGAGAGA
85851	GATGGAAAAG	TACTTTGAAA	ATGTGTGCAT	GTGAGAAAAA	TGCTAAGTGT
85901	TCTGTCTGGT	CACTTCAATG	ACAACTTTGC	TACTTTAGAA	ACTTGACTAA
85951		ACARARANIA			
86001		CAGIAGGGT.		ATAACCACCC	
86051		CAASCOCTAS		ACAGCAGACA	
86101		GUGTGTGTTA		ATTATAGCCG	
86151		AAAGTTGGTG			
86201		TOTTOGARAA			
86251		AAACAGATTT		ATCAGGAGCA	
86301 86351		AGTAATTAGT			TAAGTAATTT ATGAGAATAT
86401		AAAGGTGTTT		CCTTTTATAG	TAAGTGGTCA
86451	GTCATGACTC			ACAAGATCTT	CACTCTTAAC
86501		TGTTGTAATA			ACCATCTTCA
86551		TTTTGCCTCT			GGGTAATGCT
86601		GAGGAATAGA			CTCATTTTCT
86651		GCCTAAGCTG			
86701		TTTGACACTG			
86751		TAACTGCATA			
86801	GAAATAACTG	TAAACAGAAS	TGCCTAGTGC	ACATGCAAAG	GATTATTGGG
86851	GCTTTCTACC	CTTCAGGGAT	TAGAAGTTGA	TAGTAGGCAA	CAAGTTATAA
86901	GAAATACAGT	CAATTGTCTG	CTGACCAGGG	CTAGAGTTAA	TTGTCTCTGG
86951	AAAAAAGGAC	TTGCCTCTCT	TICTCTTCTT	CCTCCAAAAC	TTAAGACGTT
87001	TGCAGCTGAA	TCCCCAACAG	GATTTTGTTT	TCCTTTGGGA	GAGAGGAAAC
87051	AGACCAATAT	ACCCCCAAAA	CTAACCCCAT	AATTTCATTT	CAGCAGTAAA
87101	GTGAGGTCCT		CCTGCCCAAC		GTTGGGAAAC
87151		CATGCATGGG			
87201		AAATCTCATC			
87251		ATCACTCTAC			
	GGGAGAGGAG				
	GTGGAGCCTA				
	TTTGGACCAT CTGCCCCTGT				
	AAATTGCCGG				
	TGGGGAAGAA				
	CACAGAAGGC				
	CATAATTCCT				
	TGCTTGATGG				
	GAGGTGTATG				
	AGAAGACAGA				
87851	AAATGAGACC	AGGTTCGGAA	GAGGAAGAGG	GCTTGCAGAC	CTGAGTCATG
	GGGACAGTTT				
	CTTACATAAT				
88001	AAGATTTTGT	TTTAAGAAGA	TTTTGTAAAA	ACAACTGAAC	AAATGCAATC
88051	TCCTGCCAGA	GCAGGCAGCA	GCAAAGGAGA	TTAGGAATAT	AACCCCCTTG
	GAGACGTTCC				
	AGTAGGGTCA				
	TTTTTAGAAA				
	TAGGGTCAGT				
	CTGGGGTCAA				
	AAGTTACCTG				
					AGAAGGATTA
	AATGTATGTA GAGGGGAGCA				
00301	. JAGGGGAGCA	-CCIAMGGIC	CICCOMMITT	AGGAGAACTA	. ARAMICITAC

FIGURE 3, page 23 of 57

88551	ACTGACTTCT	CCCTTCAACA	GCACCTTCAG	AATCTCCTTC	ATTTTTCATA
88601	CTGTTCTTTC	AACCCTTTGA	TGAATGAGAA	ATTAGGCATT	CTTTCCCTGC
88651	AGATTTTCCC	AAACCTTCTG	· CTTTGGCCAA	TAAACATATT	TTTAGTCCCA
88701	ATCTTGCATG	CTCCTTTGGG	ACTTTTCATC	TGATAAACAT	
88751	GCTCTTGAAT	CCAATACCCT	TCTTCCCTGC	CCTCCACCCA	GAGTCTCCTT
88801	GTATCTGCTG	TTAGGCACAA	TGATGACCCC	ACCAAGGTCA	GACAATGGCT
88851	GTGGCCTCAC	CTGGACCTTG	ATGACCCACA	TAGCCTAGAG	CCCAGAGATC
88901	AGCCACTGAT		AGGGCAGTTG		
88951	CAGCCTGATT	GTTTTGACAT	GCCTGACTTC	AGGCTGCTAA	AAATGAGCTC
89001	GAGGAATCAG	ATAGGAAAAA	GAGATAGGTG	ATGCAATTTT	
89051	CCAATTTTCT	GAGTCAAGAG	TTGTTTGTTT	AACTCCAGTT	
89101	TTATCCAAAT	TTCCTGGGTG	CTTGTCCAAA		
89151	AAATTAGAAT		GACTTAGGAA	TTGGCACTTT	
89201	CCAGATGTTT	CTAATATGAG	TACTTCAACC	ACTACCCTTA	
89251	GCCTAGGACC	CTCTCTTCTG	GCAGGTGAAG	TGGAAGGAGG	
89301	GGGAGATTCT	CCACTTCAAC	TTGAGTGTCT	TGGCTTGTAT	
89351	TGGTTCTATT	TCACCAAAGG	CTTTCATCTT	CACATAAATT	TTCTTCAGCT
89401	TTAAATAATT	AGTTTTGGTA	ACCATTGGTA	TACTGGAAAG	
89451	TTGGAGTCCA	GGTGGCTTGA	GTTCAATTCT	CTGCTCTGCC	ATTTACCAGC
89501	TGTGTGACAT	TGGGCAAGTT	GCCAACCTAT	CTATGTCATT	TCCTCATGTA
89551	AAGATAATCC	CACTTCACCA	GGCCACTTTT	GAGGACCCAG	TGAAATGATG
89601	TGTAACCATT	TTAGGAACAC	TGGATCATTC	TACAGTGCAA	TTTTTTACAT
89651	CAGCTTGGAG	CCTACCATGT	AGGCATTCAA	ATCCACTGAG	TGTATGGAGC
89701	TCCGTGCACA	AATAAAAGGA	CTTCTCTTTT	CTGCCCGTGT	ACAACTTTGG
89751	TTTCCTTAAT	CAATAGAATC	CATGACAATC		GTATAAAGAT
89801	GGGACTTTCT	TCCTGTGAAG	GAGTCTGGTC	TGAACATCTT	CCAAACTCCA
89851	ACATAACTGA	TGTCATTTCT	CCACCCAACC	CCATTTGCTG	TCTCCTGACT
89901	CAATTGCTAG	AGAAGCCACT	TAAGGAAGGT	TCCTGGAGTT	AAGGCTGTGT
89951	CTGGGCCAGT	GTAGCGAGCA	GTTTTCAACA	GTCAGTCCTC	TTTATCTTCT
90001	CTTTTCCTGC	GAGCCTTTAC	TAAGCACTGC	CTCCTCCTGT	CTCCTTACTG
90051	CATCTCCTGA	TGGAATGCAC	AGGTAAATCT	CCTTGGAGAG	TACCAGCCAG
90101		CAGCCAAGGC	CACCGATCCT	CACCGCTGAG	CTCCATCTTT
	CCTTTCAAGC	TGTCCTTCCC	CTCCCCTCCC	CACCATCACC	ATAGCAACAC
	AGTGGTATAA	AAAAATGAAA	GCGCTAAGGC		AGTCTGAGTA
90251	TCAACTCTTC	CAGCATGGAG	CCGAAAACCT	AGGGAATGAC	AGCTAGAGGC
90301		TAACTGGCAG		<b>GGATAAGTCA</b>	AAGGAAGGGG
90351			AAGGGAACCA	TCACTTGCTG	AGCCTGCTGC
90401				CCCAATGTGA	ATGTTATCAT
	CTCCAGGTAA	CTGCTGAAGA		CAAAGAGGTA	AGAGATTTGG
	CCAAGGTCAC	ACAGCTATAA	GCAGTAGAAC	TAAGATTTTA	ACTCAAGTTT
	CTATGGCCCC		TGTTTCTCTC	TCCATACCAC	AGGGACAGGT
90601	GCAAGTGAGA		GAAGCACTGG	GCTTTTTGAG	CAGGCCATAT
_	AAAAATTCTG	AGCCCAGAGC	TCAACTAAAT	TATTGGAAGA	GACTGGGCCA
	AATATAAGGC	TTCTATCTAA		TGTTTCTCAA	GGACTGAGGA
90751	AAATGAAGGG			TTTCCCAGGG	TGCGTGATTA
90801	TATGGCATGG	GGGTGGGGC	CATTATGATG	CCCGGACATG	GAACTTACAC
90851	CAGTGCAGAA		TAGAAGCCCT	AAGCCAGAGA	ATGTTCAGTG
90901	TGATAAATGC	CATTATTTTT	TCCCTCATTC	ATTCAATAGA	TTTTTTTTT
90951	AGATGGAGTC			GAGTGCAGTG	
91001	AGCTCACGGT	AACCTCTGCC		AAGCAATTCT	TGTGGTCCAG
91051	CTTCCTGAGT	AGCTGGGATT		ACCACCACGC	
91101	TTTTTTTTT	TTTTTTTTT	TGTATTTTTT	AGTAGAGACA	GGGTTTCACC
31121	ATGTTGGCCA	GGCTGGTCTC	GAACTCCTGA	CCCCAAGTGA	TCCACCCACC
91201	TCCACATCCC	AAAGTGCTGG	GGTTACAGGT	GTGAGCTACC	GTGCCTAGCC
91201	TCATTCAACA	GATATTTTTA	TTAAGCATCT	GATGTGTGCT	TAACTCTGGA
01351	AATATAGGGG	CACARGAAC	AAATGCAGCT	CCTGCCCTTG	TAGAGCTTAT
31331	TAGGATAGTG	GAGAAGACAA	ATAAGGAAAC	AATTATACAA	TTGATTGATT
01461	CTTTACAACT	GTAACATGTA	CTATAAGTAC	ATAACAGAAG	AATATCACTT
91401	GCCTGATGAC	TTCAGTGAAA	GGGAAATACA	GAAGTTCTTA	CAAATCAAAG
91501	CAATCCCCTG	GGCCAATTGT	AAAGGTGATG	CCCACTTTCA	AGGTGGACAG
91501	AGACTGTGCT	AGAAGCTTAG	CCTCAACCAT	GGGTTTATAT	GATTGGTAGA
91661	CCCTGCAGAT	CCATTCCCAA	TGGTGTATCT	TCATACTAAT	CATGAAATCC
91701	ATCTAATAGC	CATACAAGTG	AGGTTTTAAA	ACCCAACAAA	CTAGACTCAA
91701	ATGAAATCTG	ATGAGGGAAT	TTATGATTTG	TTCTTCCTAC	AGCCTTTGGT
91901	ATCACTGACA	TAAAACTGAA	TGTATGTGCT	GAGGGTGCTT	GTGTCTTGGT
31061	GATAGACAAG	GTAGGTGGTC	CAGCCCATGG	TACTGGCAGC	TTAAAGTCAG
31001	CCAGCCATCA	GTGGGAAGTG	CCTGTGAATT	ATGCAGGAGT	GGGAGGGGAG
91951	GGAGTAGGCA	GIAAAGTAAT	GCATTTCTGT	GGATCCAAAG	CTTTCCAAAC
31331	TACCTGCAAG	TUAGUAAATA	TGGGGGATGT	TGTATGACTA	AGTGAGAATC
92001	AGATAATATA	ATGTGTATGG	AGCTCTTTAG	TTCTTCAGAA	AAAAATGCTG
92101	TCTAAACAAA	CHACCERCO	ATCAAAGATA	ATGATACAGT	ACCCTAATTT
92151	TAATGCTCTG	ACAMOMOCOC	GCCAGCTGTT	TCCCAGGGAT	GTGGTAAAGA
92131	TGAATGGGCA	AGATCTGGGA	AAGTGTTTTG	AAATCCTTGA	TTAAAGGCCC
92251	TCCAGGCAGA	TUTAGAATTT	LAAATGTGTT	ATATTACTGC	CACTATTGTT
92301	ATGCTTTCTT	TENTUNCUUC	MUNAATTTCAC	CATCTCCTGT	TTCAGGTGAA
92351	CGAGTCTGCC CCCTGAGATG	TORCICITAC	ACARACTECA	TGGCATTGGA	AAGGTAGCAG
	COLUMBAIG	AMIMINO	ACAAACATGT	TTTTAACCAA	GGGATCAGGA
		_			

FIGURE 3, page 24 of 57

92401 GGCCTTCCTG GCTGGCTCCT GTCAGCTGGT CATCACCTCT CTATAACTCT 92451 AGGCTTTCCC AAGCTTATTT TATTTCCATC AATAGGACAG GAATATGTAA 92501 ATGTCCTGCT TGAAATGAGT ATTGGCTACA AGCCATCTGC CTCTGAACAG 92551 AGGTGAAAAG TGGAAATCGG AGGAAGGGCA GATGTCTTTT GCAAGGGAAA 92601 CAGACTGTTT TCTGCCACTG CACTCTGCCC AGGCAAAAGA GTAAAGGAAC 92651 AGCACTCAGG AGAATTCACT GAAGCGAGGG CAGGGTGCAA AAGGAACTTG 92701 AGAAATTGGT ACTGGGACCC AAAATCAGAT TCTGGCATTT CTGGGAAAAG 92751 AAATGGGCAT GGGTGGGGGT TTTATCTGTC AATAAAAGCA TCCAGAATGG 92801 GGCTAGAAGG AAGTAAATTC AGTTGCCACC TCTGCCTACT GGACAGCCAC 92851 3GAGAACTTC TCCTTATCCA AGGTCGAGGA GCCCTCCGGA GTACATACTG 92901 ATACCATTGG TTCTCCCACA CATACCCCCA TGGAGATAAA AACAGGACCC 92951 TGGAAGCCCT GTCCGTGTTT AACCAATGGG ATTGAAACAT GGAAATGAAC 93001 TGCCCCACAA TCCACCCTGT GAGAGACCAA AGAGCAGTGT TGGATTAACA 93051 GGGAATGTTA CCCTGAAAAG GCATTCAGCT TCCACTGGGG CAGCAGGTAC 93101 AGTGCAAAGA TGATCCCACT TAAATTCCTA AGACAGGAAA TAAGGAAAGA 93151 TGTTGTGGAA ACTCAAGACC TCTCAAAGCA TACTCCTTTG TAGTTCTTCC 93201 GCAGACCAGA CCACGGAATT CAGAAAACAC CCTACCTGGT TCCAAACCAG 93251 CACCTGCCAA ACTTCTCACC CTCTTCTGAC CCTGTCCTGG GAGTTAAGAA 93301 AAAAAAAATC ACTTTATTGG TTGCTCCAGT TATAACTTAA ACAGACAGAC 93351 CATCATCAAA TTAAGTGACA TGTACGACTG CTTATTGTAT GCCAGTTACT 93401 GTGCTGTGGG GTTTTGGTTC CATTATCTCA TTTAATCCTC TCAAAAACCC 93:51 TGTTAGGTAG GTTTTATTAT TGCACTCATC TTAGATTAAG GAAACTGAGG 9350: CTCATAGAGA TTCGGTAATT TGTCAAAAGC CCTAAAACAT AATTACTGCC 93551 TCCAGATGTC TCTGATTCTA AGGCCCAGGC TCTTAATCAG TAAATGATCA 93601 AATGAATAAT GATTTTCATG GCATCTGTCA TCGGAAAGAA CAATGGAGAA 93651 TATGCTTAAC CAAAGTCATA ACCAAATAAA TGAACTTGAC AGCAGAGCCG 93701 TGATTCTAGC CAAGATGACT ATTTTCATGC ATGTTTTGAA GGCCAGGAAA 93751 AGGAGGTTAG ACTTGTTTGG GAAGGGAAAC AGGAGCTATC AAGGTGAACT 93801 TTTCCTAAGA GTAGCCCAAT AATAGTGCTC GGGAGGGAGT AATGTGTGCA 93851 AGAATAGAGT CAGGGAGACC AGCCAAGTGT GTGCCTCAGC ATCCCTAGCA 93901 CAAATCACAC ACTAAGCATT AAGATTGTCT CTGCAGTGAG AAAGGCCTGG 93951 GACCAAATTT GGGCTCCACC ACTTACTGGT ATTCATTAAT CATTCATGCA 94001 TTCATTCAAC AAATATATAT TGCGTGTGGT CTATGTGCCA GAGACTGTGC 94051 TGGGTGCTGG CAAAGAACAC AGACAAGGTT CCTGCTCTCA TGGAGCTTTT 94101 ATTCTGATGA AGGAAACAGA CCACTTACAG ATAAATAAAT AAACAAGATA 94151 AAGGGAAACA GATATGATGG AGAGTAGCTG GAGGGCCAAG CAGACCGGGC 94201 AGACAAGGTG GTGGCATGTA AGCTAAGACA TTTAAAAAGA ACCTGGTCAT 94251 GAGACTATCT GGAGAAGGAA AGCTCCAGGC AGAGGAAGCA GGTAGTGCAG 94301 AGGCCCTGAG GCAGGAATGA GGACAAGATA TTTGAGAAAA CAGAACAAAG 94351 GCAGGCATGA CCAGGCCGAG TGGGTGGTGG AAAAGTAGTA GAAGGTGAGT 94401 GGGGGAGTGG GGGCATCAAG GTCAGGCTTT GCAGGCTTGA TCAGCGTTCT 94451 CACTGTGGTT CTGGAGCCAG CAGCATCAAT GTTACCTGGG AACTTGTTAG 94501 GAATGCAAAT TCTCAGGCCC CACCCAGACC TGCTGAGTCA CAAACTCTGG 94551 GATGGGGCAC CTCATTGTGT TTTATCGAGC CCTCCAGATG ATTCCGAGTA 94601 TGCTAAAGTT TCAGAATTCC TAGGTTGGAT TATGCAGTTC AATTTTAATT 94651 TTAAATGCAA TGGGAACCTA TGAAAGATTT AAGTAGGGGA GCAGCATGTT 94701 ATAATTTCT TTAAAAATT GTTTTTAAGC ACTCCTGCTG AGGAGAGAAT 94751 GGACCATAAC AGGCTAAGAG AAATGGAAGC AGGGAGATAA ATTAGGTGGT 94801 TATTGCAAGA GGCCAGGTAA GAAGAGAAAG TGGTTTAAGT AGGGTGGTGT 94851 GGCAGAGAG ACGGTTCCAA GCAGAGGGG ACCACGCTGA CAAATAAGCG 94901 CGGGCCACTC ACGCAAGCCC AACAAGGCAG AAGGCAGAAG GCAAAAGTGA 94951 AGGCCAGAGA AAACTGGACA CCACCTTTCC AGAGCACAGT TCAAAGGCAA 95001 TGTCCTCAAA GAAGACACTC CACCCTCCTC CCATTTCCTC CCTATTGCCT 95051 AAAAATAAGA AGGATACGCG GCCTATGGCA AACCTTGGGC AGGCACGTGG 95101 GAGCTGAGCT CTTGCAAAGG GCAGATAGTT CCTCTGGTGA GAGAGAAAAG 95151 GAAGGGCCAG TGAGGAGTGA AGGAAGAGAC GAACAGAGAC CCCGAAAGGC 95201 TGAGAACGTT GTCTGGCTTC CTGAAAGGCT TAAGGGGTTA GCTCTGGAGG 95251 GTGAACTAAA AGCCCTAGTT ATATTAAACA CACACGCACA CACGCACGCA 95301 CACACATGCG CGCACACAC CACACATA CACACAGTTG AAGGAGACCT 95351 GCAGTTTCCA AAAACAAGAG TTGTATTTTT TTTGTTCATA TCATGACCCA 95401 TAACAATCTC AAAAGAGAAA CAATCTCTTG TCTTCCTTGT TTAGGCTTAG 95451 GAGAACCTGT AGTAAGTAAG CAGCAGCAGC GGAACTCAAA CTCGACTCTT 95501 CCTACTGTCA TTCTCTCTAT TACACCACAA GGCATCAGAG GACCACTAGA 95551 GTCGCCTCCC TAGGGTTAGG GTTAGGGCAA GGTAAATGAA GTGAGTCAGC 95601 AAGGGCAGGA TAGGAACCTG TCTTTATTAA CATTTTGATA TTTTGTTTAT 95651 CATGGATTTG TTGCATTAAT TGCAACTTTT AAAAATCATT GCATTAAAAT 95701 ATTATTGATC TTGATTACTG AGTTTTTAGG TGTACCCTTA AATGTTGCAC 95751 CTCTGACTTA CTAGTCTCAC CCTGATCCCT GTCCTGGATC TATGCCTGTC 95801 TGTTCTATAT CAGCCTCTTG CTTTGACCAT AAGAATAACT TCAGACCTTT 95851 AAGCATAGAG GAAATAGGAT TTCTGTCTCC CTTCCCCACC TTTGTGATAA 95901 TCTCAGCTTC TGCTTTTAAA GTCTATCTCC CAAGTAGTTT GCCTACTATG 95951 TTCCTCCCAA GGTCACTAGG TTCTGTGAAA CTAGCAGCAG GCTAGATTGT 96001 CACATTAGCA CAAAGGATCC ACTATTCCTG CAGCCGAGCT GGGACAAGCA 96051 CTTAGGCCCA CTGACTCCAA CCCTTCAATA GCCTGGGACC TACGTTGTCT 96101 CCAGGTGGTA TAAAACAAGA ATTTCCCCTT TGACTGGGAG AAAAAGGGAA 96151 GAACTCTAAA TTGGAAAACA GGTCATCTCG AATTCTCACA GGTGGAAATT 96201 TCTGACAACC CCTTTGGGAC CCACAATTCA ACACACCCA AATGGGGACA

FIGURE 3, page 25 of 57

9625	I GTAGCTAACA	TGCAACCTGT	AGGCTGTTCT	GTCATCCAGI	GCCACTGTGC
9630.	L TGCACACCAC	CAGGGGGCAG	<b>CATTCTCATT</b>	GGCTTCTATC	TOCOTOCACO
96351	CCAGTGCAGT	TGTGCAACAC	TGCAGCTTTG	СТТТАСТСТА	CTCCCTCATC
96401	l GGTTCAGTCA	AGAAAATGTC	TATAGAATCA	GCTAATCTCC	CATCCACTTA
96451	L AGTCTCTAAT	TGAAATATTT	' TCTCTGCTCA	GCCCAGGGAC	_ <u>እር</u> ር አአጥር ጥጥጥ
96201	L CCTGGATTTG	CTATTTACAA	GGATCTCTAG	AAATTATCCA	CCACAAATAT
96551	GGGCTTTCTC	AGAGCTTGAG	TGGACAGGGA	ATTARGETCO	AACCCACCC
96601	. GTTTTGACTG	CATTTGACCC	AAGTCCTGAA	GAGCCAGCTC	CTCTCTCTTC
וכססכ	. CTAATTATTA	GAAGGTTTTG	TTTGGACCCA	GTGTTTCACG	<b>中で中カ中カぐカカ中</b>
30/07	. ACAAACTTCT	CTCTTTTCTA	CTTGGATCAA	ΔΨΨΨωΨΨωΨω	サーカスススサススへ
30/21	. ATTCCCAGCA	GTGAGAGAAG	ACAAGACAGA	GAGATCCAAC	ATCTCTANAC
36801	. CCATGAATCA	GATAACCAGC	CACTTGTTCT	CTTC > CTCCT	CCCAACACAM
RCGGE	ACACTGTTAA	ATAAAATGAT	TTTATAGATT	CTTCTCACTG	<b>CCTTTCCNAC</b>
30301	AAGGGGATTT	ATCAACTTCA	GGGCACAGCA	ATC ATTT ATT	CCCACACTAC
APADI	TGGCATGCAT	ATATATATAT	ATTTACTTCT	CTTCACTTAC	AAAAAAGAGA
9/001	GAATIGGAGT	TGTGAATATT	CCTGTCTCCC	TCACCCCACC	CCCCTTCNAC
97051	TGAGTCAGGA	CAAACTTGGG	GCCCAAATGG	AGCTGTAAGT	AACTCACTCA
3 / 1 0 1	CATGCAGAGA	TGAAACCTTC	ACAGACCCAC	TGATATGGAG	CTTCNNCNTT
97151	AAATTCCCCT	TTGAGAATAA	CTGGGTAACA	CTCATACAGA	CACTACTTTC
9/201	AAGAAGGCCA	GATCCTCCCT	CTAATGTATA	GTGCAACGTT	CCTAACCCTC
9/251	AGCCCACTCC	GTCATACCCC	CACTCACATG	AATACACACA	TARCCRCTAR
9/301	TATAAAGCAC	TTCCCACCAT	AGGGCAGCAA	AGAAGGAGGG	<b>ል ል አጥር ጥጥጥ አጥ</b>
3/351	TATGGAAGAG	TGGAAGGAAG	GAAGGGAAGG	GAAGGGAAGG	GAACCCTAAC
9/401	AGGAAGAATT	CTCAGGGTGA	GCAGAGGAAT	GACATGTTTG	CCCCATAACC
97451	AAGATAATTG	AAGTGCAGAG	TTTGTATGGA	AAAATTTGAA	AATATCACCT
97501	GGCAGGCCAG	GCATGGTAGC	TCATGCCTGT	AATCCCAGCA	CTTTCCCACC
97551	CCAAAGCAGG	CGGATCACCT	GAGGTCACGA	GTTTGAGACT	ACCCCCCCA
97601	ACATGGCAAA	ACCCCATCTC	GACTAAAAAT	ACAAAAATTA	CCTCCCTTTTA
97651	GTGGCGCATG	CCTGTAATCC	CAGCTACTCG	GGAGGCTGAG	CCACCACAAM
97701	CATTTGAGCC	TGGGAGGCAA	AGGTTGCAGT	GAGTCGAGAT	CARCONACMA
97751	CACTTCAGCC	TGGGTGAGAG	AGCTTTCTTT	TTTTTCTCTC	CAIGCIACIA
97801	AAAAGTTCAG	GTTGCAGAGA	TGGATGGATG	GATGGATGGA	TCCATCCATC
97851	GACGGATAGA	TAGACATTAC	AGAGAGTTTC	CAATTCTTAG	CAMCAAMMCC
97901	AATCCTTAAG	TCTTTATTCT	GTAAGAAAGG	AAGGGGAGAA	TANAMETER
97951	TGATTTTAAA	ATATTTTCTA	CCCTGTAGAG	CTACCCTACA	ACCCATCAAA
98001	ACCTTAAAAA	AAAAGGCATC	TACTTTADAD	GAATAATGTC	TARRAM
98051	GAAATTCCCT	CTTTTTGCCC	TGACCTTTGG	GAAACAGAGT	CACHCAHCCH
98101	TTTGAGGTTT	TTGGCACTGC	CTTCCCTCTC	ATCATATCCT	GAGTGATCCT
98151	TCCATAATCA	TGCAGTTACC	TCAGATGTCC	CTTTCCCTCT.	ACCOLCIAGE
98201	AACACGCTCT	CCAGGCACTG	GGAAAGTGGG	TAATTAGGAA	AGCCACAGGT
98251	TACCCATGGG	CTGTGATGCC	CAGTTATAAA	CCCAGACATT	MCACARGUAG
98301	CAGAATGAGC	ATCAAGTCCT	CAAATGGGTC	TACATCCATA	AACAMCOCCA
98351	GCAGTCAGCT	CTTTACTGTC	AGTAGAGACA	AAATGTTCCT	AACAIGICCA
98401	TAGGGGAAGC	CACATCCTCA	GTAGGGGGGG	TCTGATGAGT	CCACCTTTCCC
98451	ACAGGTATGT	AGAAGCTGCA	TGCAGCAGAG	GGCTCAAAGG	ACCOMMON
98501	ATAGATACCA	AAGCAAAAGG	GGAGTCTGTG	CACGTTCTCA	CACCACAC
98551	GAAACACTCT	TTTTGTTCAC	AAAATACATC	GTGTAGGGTA	CAUGUACCCC
98601	ATCATTTAGC	TCAGGTTCCT	CCCCCCATAA	AATAAATAAG	COMMONANT
98651	TAGTTGTCTG	TTGCTGTGTA	CCD D D TTCTC	AGAAACGTAG	CCTTCCATAT
98701	CAATACCCAT	TTATTATCTC	GCAACTTCTC	TATCTCAGAA	AGGCTTAAAG
98751	GCTTGACTGG	GTTCTCTGTC	CARCTTCTCC	TGAGACTGAA	GTCCAGGCAG
98801	TGGCCAGGCT	GGGATCTTAT	CTGCACCCTC	TGAGGACATA	ATCAAGGTGT
98851	ACCTTATTCA	GGCCATCAGC	AGAATCCCCT	CTCTTGTGGC	TACGCTTCCA
98901	AGGTCCCCGT	TTCCTTCCTC	GCTGTCATCC	AGGGACCACT	TTGAGGTTGG
98951	ACAGGCTGCC	TATGTTCCTA	TTCACAACAC	ACCGTTCATC	TTTGCACCT
99001	AGCAGCATGT	AGAATCTTTC	TTGTGGGGTCG	TATCTTTCTG	COMMUNICATION
99051	CTTCTTTAGC	CAGAGAAAGT	TCTTTCCTTT	TAAGCGTTCA	TCCCTT TCCCTT
99101	TCAGGCCCAC	CTGGATAATG	TCCCTATTT	AAAGGTAACT	CECAMA CCC
99151	ATAACATTTC	AGGAGTGATA	ACAGCACATT	TACAGGTTACT	AACCAMMCCC
99201	GCAGAACATC	TTTGGGGGAA	CATTTTACAA	ACTCTGCCTC	AAGGATTGGG
99251	CATAATCCTT	TTAAAAACCA	AATCTTCAAC	CCTTTTTTTC	CCCACTCACC
99301	TTTTGAATAA	GCACATTTAT	ACCTA ACTTC	ATCAGACACC	CLAAAGGCCT
99351	AAACACTAGC	ATGTGGCAAA	ATAGGCTGTA	AATCAATCAG	AACTTTGAGC
99401	TCCCACCACA	ATCTTTCTCA	ATCACATTCC	GAGAATCTGA	AACTATTCTT
99451	GGTATACCAG	AGCAGACTCC	TACCACATIGG	CAAGAGCTGA	CACTGTCAGT
99501	TTTAGTAATT	GTGGACATTC	CTTCTTNAAA	TATTAGTAGC	CTGTTAAATG
99551	CTATACTCAC	AGTATTTTTC	CCATCCAAAC	CAACCGTTCC	LIGAAATTGA
99601	TTCTCTTTAT	TCCTGGGAAC	CTCCTTTAM	AGCTCACCAC	MAATCAGGGT
99651	CCTTTAGGGG	TCATTACTTC	ACCTOCTOTAL	GCATGCAGGA	LUCCTGTAGT
99701	TGGCCTTTTT	TATGCATCCA	CDTCTCTGTA	TTTTTTAATA	ATCCTCTCCA
99751	GGTGATCACT	CTCTTATAAC	CTACTOCATO	TTTTTTAATA TCCCTGATGG	CCAGGAATGG
99801	GGTAGAGTTG	AAACCCACCT	CTUGITORIC	TCCCTGATGG	AATGGTATGT
99851	AGCAGCACTT	GTGACACCCC	CACAACCARR	TGGAGTAAGT	TICCTTTGGA
99901	CCAGGAGACA	<b>でこりれいれいししし</b>	CCATCCACT	TGGAGTAAGT ATCAATAGTT	AGCATTTCCT
99951	TCTTTAGAGG	CACACTCTCT	CAATCCACAA	ATCAATAGTT	AGATGCAAAA
100001	AGTTATAACC	TTACCCAAC	CTCTT > CTC	CCCAGCTCTG	CCACTTATTT
100051	ATGTGTGGGA	ATECCCAMEN	AAATAGGGG	TTCTGGTCCT TACCTCATAG	CTGGTTCTTC
		ALGOGGATAA	AAA FAGCACC	TACCTCATAG	GTTATTATGA
		T-1	TOT TO T		

FIGURE 3, page 26 of 57

			•		
	ATATTAAATG				
	TACCTATCTA				
	AATGCTTAGA				
	AAGTTAATTT				
	ACTCTGGATG CTAATTAGTC				
	CATCCTTGAC				
	CTTTAATAGA				
100931	AGTAGCATCC	ACACACCCTC	CCCCCTTCTT	DADARGCAG	CCTCTCACCC
	CCCACCCCAG				
	GTGATTCATG				
	ATGACACACT				
	CATGGAACAG				
	ATCAGTGTTG				
	TTCAAAGAGC				
100851	ACCGGAAAAA	TAAAGGAAAC	TACAAGAAGA	ACCCAGCTAA	GAGATGTGAG
	GCTTCTGAAA				
100951	CCAGCTGCCC	CAGGTCAAGG	AAGCTCTGTG	AGTGTTAGCT	GACCCGGAGC
101001	AGCAAGGATA	CATTCAGAAG	TGATGAAAGG	GAACGCTTCT	TGACAGGGTA
	AAGAGTCATT				
101101	CCAGCCTGTA	CTCAGAGATT	ATTTCTGGCA	TGGGAGGGCC	GAAGGGTTAG
	GAGGCCACCT				
	CCTGTGCTGC				
	CTGCAGCTTG				
	GCTATAGGTG				
	AATAGCTGTG				
	CTCATCTTTA				
	ACCTATAATC CCAAGAGTTT				
	AAAATTAGCC				
	GGCTGAGGTG				
	CCAGATTGTG				
	CAAAAATAAA				
	AGAATGAAAT				
	CTTTGGGAGG				
	AGCCTGGGCA				
	AAATTAGCTG				
101951	GCTGAGGCAG	GAGGACGGCC	TGAGCACAGG	AGTTGAGGCT	GCAGTGAGTC
	ATGATCACAC				
	CAATAAATAA				
	ATGGTGGTGA				
	CCTGTATAAA				
	AAATGCTGCT				
	GAACCCTTCC				
	TCCTCGATGC				
	GTATTTCCTA				
	GGTTTCCGCA GGCACCTCCC				
102501			GCTCTAGGAG		
102551			CTGCCACTGT		
102601			GAAAACCCTC		
	CCCTGGGTTG				
102701	GTTAAGTAAA	CTGGGGATCT	AGGTTTGATG	ATACTGGGTC	TGCAGCTTCT
	TTGTCCCACT				
	TTTTTGTCTC				
	TTTACGAAGT				
	TTATGTTTGA				
	TACTGTAGCA				
103001	AGGGTGGGGT	CTGGTTAGTC	CTTGAATTGG	AGACTGCCTG	GAGATACTGG
102101	ATGCTGCAAG	CTTTTGAAAA	AAGACAAGTT	CTCTGTACTT	GCAGAGCTTA
	CATCCAGTAA GGTGGAGCCA				
	AGAGGTGATA				
	CCCGGTGAAG				
	GGTGGGAAAG				
	ACATAAGCAA				
	ACTGGATCAT				
	ACAATTACAG				
103501	CCTGGCTATA	ATGTGGGGAA	GGGATTGAAG	AAAGAGGGCA	AAGGCAGGAA
103551	CAGGAAAATC	TCTTAGGAGG	CTACTGCAAA	GCCCAAGGGA	GAGGTGATGG
103601	TGTTTTGTTG	TTGTTGTTGT	TTGTTTTGTT	TTGCTTTGAG	AAGGAGTCTC
103651	ACTCTGTCGC	CCAGGCTGGA	GTGCAATGGC	ACAATCTCGG	CTCACTGCAA
103701	CCTCCGCCTC	TTGGGTTCAA	GCAATTCTCC	TGCCTCAGTC	TCCCAAGTAG
103751	CTGGGATTAC	AGGCATGCAC	CACCATGGCT	GGCTAATTTT	TGTATTTTTA
103801	GTAGAGACAG	AGTTTCCCCA	TGTTGGTCAG	GCTGGTCTTG	AGCTCCTGAC
103001	CTCAAGCGAT TGAGGCACCG	CCCTCCCC	ATCATCCT	AAGCACTGGG	ATTACAGGTG
エハフコロア	TONOGCHCCO	COCTOCCCAN	WIGHT GOLCL.	TTTGATCTGG	GICTIAAAGG

FIGURE 3, page 27 of 57

103951		GGGGGTAGTA	AATTAACTGT	GCTGGGGAAG	AGAGGGAGG
104001	CTGAGAGTGA	GGAAAGAATG	AGGGGTGAT1	CCAGGTTTAG	GAAAACTGG
104051	. CAATTTGTTA	GATGATGGT	CCATTGACAG	AAATGGGAAA	GAACAAGTT
104101	. GGAAAGAAAA	CTCAAGATCT	GGCTGGTGAC	TTGTATTAAA	CTTABACCC
104151	CATTTGTGAC	TTGAGCAGAA	GTAAGGACTI	TCTCCAGTGT	TCAAGAGCT
104201	GAAGGGATTT	' TTCTAGCCTC	CAGGCAAGGT	' AATACCATAA	GTCCCAACAC
104251	TGATGCCCTC	CCTGGGAATG	ATCTCAATGG	GAGAATCCTA	TACCCTGCCT
104301	CCTCCATTCA	. TTCCTTGCTC	TGATGGTGGT	TCTGGCTGGC	TAACCTAACT
104351	TACTCTTGCC	ACTAGTTAAC	GCCTGTCCTT	ATTTCTCTTG	TCCCCACCTZ
104401	AGATGTCAAT	CAAAACAGCA	CGAGCCATGC	TATGTCACAT	GACATGTTGT
104451	CTGTCCAGCC	CAGAGCTTGT	'- TGCTGATGGG	GGCACAGACT	AGATTTTCAC
104501	AGAAATCTCT	CTGTTACCAC	CCTTAACATT	CCAACCCCCT	CTAATAGCCC
104551	ATTTAGGATT	TATCATACTG	TTTCATCCAA	ACCTTTCATG	ACCTGATTTC
104601	TATTTCCAGC	TTCAACCACC	CCTTGGGTCA	CCACCTGTAC	TTATTGAGTT
104651		TCTGAATTAA	TGACTGAAGA	TGATAAGCTT	CCCTTACATA
104701	TGACTCTCAA	ACCACCAAAC	TGGGATTGTT	GTTACTCTTA	GTGATAATGG
104751	CITATITA	TGAAACTTTT	AATAGGGAAC	ACAAACCCTG	CCCAGAAATT
104851		ATTTCATTTA	AGAACATCAC	AAAGTAGGTG	CTATTATTTG
104901	ACCTACTOR	-TGAGACTTGA	AGAACTTTAG	AGCATTGCCC	AAGGTCACCC
104951		AACCA ACCCC	GGGATTTGAA	TCCAGCTCAT	CTGTCTCCAT
105001	GCCACGGGCT	TTCTACCOTA	AGAGCATCAT	GGCCTTTCAC	AAGTTGAAGA
105051	GTGTGGCAAG	TCATCACCCTA	TTCCA CTTCA	CTTTTCCATG	ACTGGGGTGG
105101	CAGGTGTCTT	CCTAACTCCT	CEECCARRO	TGTGGTGGGG CTTCAGGAGC	TGGCAGGGAC
105151	ATCTGATTCT	GCAGGATCAA	CAATATCA	ACTGCAGGCT	AAAGGACCAG
105201	CCAAAGCTCT	AATGGTGACT	TCCCCAACCE	CAGGAGGGCA	CTGTAGACAT
105251	ACCCATTTAG	AATGTAAACA	TTCCTATTT	ATAAAAAAAGA	GGGAGGTTGT
105301	ACTGAAGGCC	TCAGTCTCCT	CCDACADAGC	CAGGCTGTGG	AAAAAAGGAG
105351	TCTCAAAGGG	TGCAGGCCCA	TGGCCACTGC	CCAGGGCTCC	TCCTCACCCC
105401	TCCTCACTCC	CACAACTGAG	GGGAGACCCA	GTTCCACACC	CACCCACCEA
105451	GCAGTGTCTC	ACACCCACCG	GGAGAGGTCT	AAACATCTTC	CACCCACCIA
105501	GGTCCCAAAA	TGTCCCTGCA	GTAAGCAACC	ATCTGGAGAG	GCCCAGGTCT
105551	ACATCTGTTT	TTAAAGCTCC	AATAAATAAA	TAAATGAAGG	AACAAAAAAA
105601	GAAGAAGAAA	TGCAGAACAG	GGTGACTAAA	ATTGGCATGT	ATTTTTAAAT
105651	GTTTATATTA	ACAAACTAAC	ACCTTTTAAC	ATGAAAAGCA	ATATAATTCT
105701	GCTAGCCACA	AAATCATCGT	AGGACTGAGA	AAGGAATCGT	GATTCTGAGA
105751	GCCCTAGAGT	TAATGTGATC	CAGCTGGCTC	ATCCCTGTGA	CTGCAGAAGC
102801	CTGTTTGGAG	ATAGTGTCAG	TAGCTTTTCA	GGCCCTCTGT	GAATTGCCAG
105851	AATGTGTGAC	ATGAGCCAAA	TTTCCCCCCA	GCATCCCCCC	CGCCGCCACC
105901	ACCACCCCC	ACCCAACCCT	CCCGCCGGCT	CCCATAGAAT	AGTCACTGCC
105951	ATACAGAAAA	AGAGAAGTTC	TACTATTTCT	GGGCAAGATT	TCCACAAACC
106001	AGTTTGTCCC	TTTCTGCTTT	CATGAAATAA	ACCATTTGGA	TCAACGTCAG
106051	CTGATTGCAA	AAATTTTCCC	TTGTCTCAAA	AGCAAGACTG	ATAAGGAAGC
106101	AAACATGGGA	GGACCTTAGT	GGCCGAGCCT	TTATGTGTAT	GTTATTTCAT
106151	TGCTCTCATA	ACTGCCCTGG	GATGCTGTAA	GCATGATTCA	TCCTGTTTGT
106251	TTATCAGTTA	AATTATGTAT	CCAAGATTAC	ACAGCCTATC	CAGGATTAGA
106301	ACTCAGAGCC	CTCGGCTGTG	AAGCTTGAGC	TCTTTCTTTT	CAGTCTTCAA
106351	ATATGATCAT CTGGAGGGGT	CCRCRCCCA	AGCACAAAGC	CCAGGAGGAG	CCCAGTGAGG
106401		CARCTEGCAG	CCACTCTCCT	CCGTGCCCCT	GTGGTGTTGG
106451	GGCAAACTTG TTTTGTCTGC	TCCCTCACTO	AATCTTTTAA	CTGTTTCCTT	CTCTTCCCGT
106501	TTATTTTTCC	ATCCACACCA	AAACAACACT	CTACTCCTTG	CTTATGATAC
	TTATTTTTCC AGGCATATGA	GAAAACAAG	AMMCMMIICA	CATCAAGGTA	ATTGATGATG
106601	ATGGTGGATA	TGAGTTTTCA	CANACCTCA	CTACCCTCTC	GGGCCCCCGC
106651	ACACTAACAT	TCTTCTCTCC	TCTTCTCTTT	CTTCCTCTCC	AACCCAMMMC
106701	TCTCCTCCTC	CTCTTGTCTT	CCACCTCTCT	GCTTCCCCTTT	CCCETCTCTC
106751	CTCTCTTGCT	CTCTCTCCTG	CTCTCTTTTC	ACTCCTCCCT	CTCCTCTCTC
TORROT	CTCTCTCTGC	CCCCAGCTCT	GTCCTAACAC	CTGCCAGCCT	GACACATICCC
100821	ATCCATACGA	GGGATGCTCA	AGACCGATGG	TAATTGTTCT	GGGATAAGGA
106901	AATGAGTATG	GGGAAAGAAA	GAGCCAAAAT	CCTCCACTAT	CATCTCCCCC
106921	TCTTGGCTTC	TCCAGAATGG	CTGGGCATAA	AGGGGGGGAAA	AGGGACCACA
10,001	TAGCCCAGCA	CCAGACAGAA	GAGCAGCACT	GAGAAACAGG	CTTTCACCAC
10,021	AAATTTCCAT	GGGGCAGTTA	TTCTCAGGGC	TAAACTTAGA	GTCCCACCAA
10/101	GTTGAGAATC	AATGTATTTG	GATTACAGTT	CATTCCCCCTC	CCAAAACCAC
10/121	GCTTTAGGAG	CCACCTTATC	TGCCATGTTG	CTACTATCAA	CACTTCTTTC
10/201	TCCTCCTGAC	CTTGAGGAAG	CTGAAAGTAC	AGGTTTGAGT	<b>ፕሮሮ</b> ልፎልሞርሞክ
10/521	GGTCAAATAT	CCATTTGTCT	TCCTATGTTT	ΤΤΟΟΤΑΤΤΑΑ	CAACACCCAC
TO 130T	GTGTGGAGGC	AGAGAGTTAG	AATAGTGGTG	GAGATCATCC	TCACCCAAAT
10/321	GGAAGCTTCC	CCAAGAGGTC	CATGGGGCTT	CTCAGAGTGG	<b>Δ</b> ΨርርδδΨርΨΨ
101401	TGCCTTCAAC	TTCAATGACC	CCATACATCC	CATGGCCTCC	AATACACAAC
10/451	TCAAGAAGTC	CTTTCCTGAA	TAGATCATAC	TGTGGAGCAG	GGAGCTGCCA
10/201	GTACTGAGGG	CAATGTTCCT	TCCCCTTCCA	ACCTCTCCCT	CATCCCCTCC
TO 122T	AGTACATGCC	TGTTGTCACA	GAGCACCCCA	ATCCCATCCC	ACAGCAGAGT
10,001	TCCTGCAGCA	GAGAAACAGG	CTCACACCTT	GTAGACACCC	CTGGGGTCCC
TO 102T	ATATCTAGGG	CCAACAGAAA	TATTCCCAAA	AAAATGCCTC	<b>でからなぐな かかぐか</b>
101/01	ATGAGCTTTC	TCTTTTGTCC	GCTGAGCAAG	CTATAAAAAC	ATCTC A A A A C
~0112I	AAGTACCCAA	MAAGGTAATA	AAAATGTACA	GTCGTGCATC	ACTTAGCAAT
			~~~~		

FIGURE 3, page 28 of 57

					•
		TCTGAGGAAG			
		GTGTACTTTC			
		GTGGGCCTAT GAATATTGCA			
		CTAGACATAG			
	AGCCTTATGG				
	TTTTATGTAG	CATGTGACTG			
	CAGCAAGTCC		AAGCCCCTTT		
108201	AAATGGCAAA	GCCGAGCACG	CCCACAGAAG	GTAGCAGGAA	CATCAGAGGA
108251	TCTGAAGAGA	CATTTAGGTA	AATGCTCTTT	ACCCTTTAGA	GCATTTAGTT
		CCCTCCCCA			
		AAGCAGAAAA			
	TAGCCACCAT		TGAAGGACCA		
		CACGGCAACC			
	AAGATGTTAT		GATACCCAGA		
	ACCTTGTGCC		GATAATTCAC		
		CCCTAAGTGA			
		ATTTTCTTCA			
		TCCTATGTAA			
		AAGCCCTATG			
		TCTTCTCTCT			
		CTCTCTTCTC			
		TTAAAATTTT			
		AAGGGACACA CCCTGGGTTA			
109101		GCCACTGTTA			
109151		CTACTAACTG			
109201		CTTCTGCCTT			
109251	CCTTTGCCCA	CCTTTCCTTG	CCTCCTGGCT	CCCTGCCCCC	TCACCCGTAA
109301	GAACAACTAT	GACCAAGAAG	ACAAGAAAAA	CTAAGACCAT	TTATTACCTG
		AATCCACCAT			
		TGCCAGGAAT			
109451		CCAATCTACC			
109501		AAAACCATAA GAATTTCTTC			•
		CAGGTGTGAG			
		TTAAAACAAA			
		TGTCTCCCTC			
	TGACCAATGG		TCTTTTCCCT		
109801	CTGTTTCGGT	ATCATCTCTG	CCTTCTTAGC	CTTAGCTTAT	TCCAAATTCC
109851		CCTTCTGGGC			
	AATGAGTTAT	TTCCCTGTTT	TGCTACAATT		
	GGGCCCCTGC TGAAAATTCC	ACTCTCCCCC	GGGAGGAGAA		
	CGATGCATGC		TCCTGCCTTC		•
		GATTGAATGT			
	GGTTGCAATG		AGTTTTCTGC		
		CAGGTTATTC			
110251	CCTTCGAGGA	GGTAAGTGTA	TTTTCTGGCT	GTTTCACAGT	TGGGCAGACC
		AAAGTGTACC			
		GAGTTGACTT		•	
		CTCCTCTGGG			
		GGGTGCCTCA ATTGAGAGTT			
		TAGCCAGGGG			
		ACTTGTGGAT			
		ATCCCTGACA			
110701	CCTTTGCTTG	GCATCTGCAA	GAGAAAGTAC	CGCCCAGATC	CCAAGATAGC
		CACTAGAGAA			
		AATACCTGCC			
		GTAGAAGTAA			
		TACTTTGAAT			
		AGGGTGCTCC AGATATGCTG			
		TTTGTCAAAA			
111101	TCATGTGACA	AAGAGCAGCA	TAAAACTTTC	CACACGAGGA	CAGAGCTAAG
		ACAACATTCC			
111201	GATTGGTCAT	TTCTCATTGT	CTGCTGGGGA	CTCTCCTGCA	GAGCTGACCA
		TGCGCTGGTT			
		CTTCACTGCT			
		CTGACTGCCA			
		TTCCCACATT			
		TCTCAGAGGT GCATTTATTC			
		CCAATTGTAA			
		CAAACGAAAG			

FIGURE 3, page 29 of 57

111651	CAGTGGTTCC	AGACAAGAGG	AAGAGATTGG	AAGTCCATAC	ATGCCTTTAT
111701	TCCACCAGTA	AAAAGGCTCT	TCTCTTATGC	CTCCCTTAAA	ACCTCTACCA
111751	ACAGCAGGAC	AGAGAGTGAC	CCAAGATAAG	TCTTCAAGAG	ACCTAACCAA
111801	ATGCAAATGT	CTTTGGCTAA	TCCCCATTTA	AGGACATCTT	CCTCTTTTCC
111821	ACAGATTCTT	TGCCCAAGGA	AATGTCAGCA	ATGCCCTCGT	GGAGGGAGTA
111901	GGTGAGAAGA	CAAGGATTTC	AGCAAGCTAT	CTGTGTGGTG	TGCCCCCAGA
111951	TCTCCCCAGT	GACCGAGATG	CCAAGATGAA	GAGTGCCAAG	AAGAAATTGG
112001	TCAATTTTCC	AGCTGCCTAT	TTTATTGTCT	<b>ATGTTTTCTA</b>	GGCGGTTAAT
112051	TTCCAGTTTC	TTCAGTACTT	CCCGTATTTT	GACATTAGAC	CATAAGGTGA
112101	AAGGTCATAA	AACCTGATTG	TCTAGACTCA	GAAGCAAATG	GABACCCATC
112151	CAAATTTCCA	GAATTCCCTG	CTGTTCTCAG	AGTGAGAAAC	AGAACAGTGG
112201	AAATTGCTTT	TCATTATCAC	TACTGCATGG	GAGAGTCTGA	AACATTCAGA
112251	ATGGCATAGT	CTTTGCATGG	TCAAAATGAC	AATTGCATTA	AAAAAATCAC
112301	AGACTGGATT	TGAAATAGGA	GACTCTATTT	TTGGCAAACA	AAACAGACTT
112351	CAGAGTTGAG	ATTAAAAGCT	CTGGATGAGC	TGGGGGATGG	AAAAAAGGGA
112401	AGGAAAAAAG	GGAGACTGAA	TAGGAAACAC	AGTTGCTCTG	GAGTCTAGAA
112451	GTGGACTTCC	GAGAGCAACA	CTGAGCAACA	TAATCAAGAC	TGTTGGGCCCT
112501	GGGCCTGGAC	ATTGGAAGCC	TTCGGATAGA	AAGGAAAGCT	CTCTGTCTCT
112551	CTCTCTCTCT	CTGAAGAATG	GGGCCTGTTT	GGTCCTCCTT	TTTCGACAAC
112601	CGTGGGCTCA	TCTTGACAAG	CTGCCCAGAT	GCTTCCTAAT	TACTCACAGT
112651	CCTATGCTCT	TTCCAGCTTG	TCCCTGGGGT	GTCTGAGCAG	GAATAAATGA
112701	CTCTCACCTG	ACCCAGGGGA	TCAATACAGG	GGAAAGTTCA	CCTCCACCTT
112751	CTCTCATGAG	CAGCAGCAGG	AAAAACACCC	TCGAGGTATT	GTGTCAGTCA
112801	AAGCTGGCCT	ACCCAGGTCT	TGCTGACCCA	TCTATAACTC	CTGAGCAGAA
112851	AGTCTTGGAT	TCATGGAGAC	AATGACCAGA	CARTGATECA	ATTCCAGCCA
112901	ACTGCAGGCC	TTCTCACTAC	TCTAGGGATG	GGCCACATCT	TCCCTCCCAM
112951	GTATGAGTGA	AAACCAGGGC	ATCAGGGACC	TTTCTCCAAC	ACCTCCCTTO
113001	GTCTGACCCA	CCTGTGTTCA	TTTATCTCCT	GGGATCTCTC	ATCTCCCCTG
113051	GAACTTGGGG	GAAGCTCTTC	CACGCAAACT	CCCCCAACCA	CCACAAMAAA
113101	CAAGCTCTTG	CCTATCTATC	TATCTATCTA	TCTATCTATC	TATCTATCTA
113151	TCTATCTACC	TATCTGCCTA	TCTATATCTA	TCTATCTATC	TETRETACTOR
113201	AAAGCCATTG	ATCCATTAAC	CTTTGGAATT	CTACATGGGA	CATACCTAAA
113251	AAAGTGAACT	GCCTTGTTTA	TGTATCATGC	AGACTCTGGA	TCCACATATA
11,3301	TCTCAGTGGC	TGTGAATATA	GGATGATTGA	TCACAGGCCT	CACTECATA
113351	CCTACAGATT	CTTAGGAAAA	AAATTGATTC	ACAGACATGT	CCCCCCTCCT
113401	TCCCCCACAA	CACACACTCC	TTCCTCAGCA	ATCTCTATCA	GTCACCAACT
113451	ACACGTTGAA	TATGTGGCAA	GCTCTTCCCA	GACCTTTATC	TENENCECAN
113501	GGAGTGAGGG	GCTGTACTAA	GATATCATAG	AAATGAAAAT	GTGGTGTGTC
113221	ACAAGTTTCC	TTAATTCTTA	GATCTTAAAC	TCTAAGAGGG	TTCAGCATAA
113601	GTACAAATTC	AAGGGCTAGA	GACAACCTGT	ATTGGGTGTG	<b>ጥርጥጥጥል አርጥ</b> ር
113651	AGTTTCCCAA	TCCACATAGG	GACCTTGCAT	TTGTCATCTC	TCATCTATCT
113701	ATAGCTGTTG	GTATGACAGT	TTCTCTGTTC	CAGAATACCT	GAACTCTGAC
113751	TTAGCCTGTC	CTTTCTGAAA	CAGAAAAATC	ACCCAACCAG	AGATCTATGA
113801	GATCTATGGA	AAAGACAGTT	GCCAAAATAG	ACAGCAAACA	GCCAAACTTA
113851	ATTGAACACT	ACCACATGCA	GGGACTTTGC	TAAGCAGAGG	TGATACAAAA
113901	TGGGAGGAGC	CCATAGCCCT	AACTTCCAGG	ATATATCTAC	GGTAAAGACA
113951	AACCATTCAA	GGAAAACATT	CTGCAGGACT	TACCTTTTTG	CTAAGTCATT
114001	CTTTTAGGGG	AAATCAAAGT	TCTAGTCAAC	GTGGCAGCTA	GGAAGGCATT
114051	TGTGGTGATG	GAAACCTTAT	GAGCACTGAG	AAGCTGAGCA	TGAGTTCAGC
114101	TAAGTCGTTA	GGGATGGAAG	ACATAGACCT	GGGCACTGTT	CCACTCTTGC
114151	ACAATGCTAC	CCATTTCCTT	GAGCTCCCAT	TCAAGCCCCA	<b>TCCTC ATTT</b>
114201	TGCCACTCAT	AAGTTAGCTA	CTCTGGCAGG	GTTGCAACTT	ACACAGTTTT
114251	CATGATAACT	GGATTCTCAC	TCCTTTTTTT	ACAGAATGGA	TGTGATAACC
114301	TGGTATCCTA	CACAGTCATG	AGTGACCAAC	CTACCCATTT	ららずずしてしてひず
114351	CCTCATTCCT	CCATTCCTAG	CCCTAGGGTA	GCCGGGAAAG	CATAGGAGCA
114401	AATGCCCTTA	CCAGGGCCCT	GGTGCTCAGC	AGCCTCTCCG	GCTGCTCACA
114451	CCTCTTGCTG	CTGCTCTGTG	CATGCTCCAA	AGGCTGCTTT	TTCCCTATCC
114501	CTGCTGAGCT	CTCACCTACT	AAGCTCTCTG	CTTTCCTTAT	GCTGCCAGCA
114551	ACCACAAAAC	CTGGTGATAC	TTTCAAGATG	GGACATTAAT	<b>CCTCTTTCCT</b>
114601	TTTCTTTCTT	CCATTTTTCT	GGTATCCATT	TGCAAACAGC	GCTCCTGTTA
114651	TCTCCAGGTA	AGAGGTGTCT	TGTCCCCCTC	TTTTCTTTCC	ACTTCTTGCC
114/01	AGTGCCATTA	TTTGGTTTAA	GACCAATGTC	CTTTGATTTA	TTGAATAAGA
114/51	ACTGCAGGCT	CAAGTTAACC	TGACAATTTC	TCCCAAGGAC	TGGGAGATTT
114801	ATTTTCCCAC	ATGAAGCAAT	TATGAGAAAG	CAATTGTGAG	GAAGGCAATT
114851	CCTTGAGCAT	CACTTCTGTC	TGGGGACGTG	GGTTAAGGCA	TAGCTGATCC
114901	TCTCTGGGAC	CAGGAAGAGA	AATTAAGCTT	AACAAGGAGA	TECTECETE
114951	TAGACTTCTC	CTGAGTCTTA	ATTCATCTGC	CATCTCATCT	TGTGGGGGAA
112001	GAGACAGTGA	GATTCAGAGC	TGGAATCTCC	ΤΆΔΤΆΤΑΛΤΉ	GTGACACGAT
TT202T	TTGAAAAAAA	AATACTTTAA	TCCCAAGGGA	TCCAGGAAAT	DACCADACCT
TIDIOI	GTTGTGAGAA	TAGGAAATGC	AATTTTTAAA	GAATCTGGAA	ጥጥጥከልሮሮክርጥ
TIDIDI	CCTGGAGATC	TTCCATCTCA	TCACAGCTGA	GACTTAAATT	CCTACAATTT
115201	TGGTTCATTT	GTCATTGACC	CTTAAAGTCC	TATCTCCCCT	CARCARCARC
115251	AATTAGGATG	GGGGATTGGG	GCAGTGTTCT	GGCTGGDDDT	ስጥ ስ ስ ስጥጥጥ <b>ስ</b>
112301	GAGAATTTAT	TTTGAAGAGA	TTCTCATGCA	GAATCTAGGT	CCTATACACC
112401	ACGTACACCT	ACTITGAGAG	TATGCTTGCA	TGAGTGGAAA	CCAATCATAA
115451	ACAACATTCA	ACTICATGAG	CAGATATGAA	AGCATTTTCA	GCATATCTAG
113431	CAATACTATA	ACTUITITGTG	CAAGCAGAGT	GCCTACACA	AGACAGTTTC
		דיד	OTTO		

FIGURE 3, page 30 of 57

		AAAAGAACGT			
		AAGGCCACTT TCACCTACAG			
	ATATTCAAAG	GGACATACGC			
	TGTAGGTAAC	TCCTACATTT			
		CTTTTTGCTT			
	CTCAACCCTC				
115851	TGGGTTTTTC			CTTGCCTGTG	
		GTTCCAACTC			
		GATTGTGATT			
		TCAATAGAAG			
116051	CCTAGCTCTT	TCAATCTGTA	AGCCTTTAAT	TTAGGAGCGC	TGATTAGCCT
116101	TTCAATTCGT	TGGAAATCTC	AAATACTGGT	TTTAATTTTC	CTAGGTGGAC
116151	AGAGACAGAG	GGAATATGTT	CATTCTGAGC	TAACCACCCC	CCCACCCCA
		CCTTGCAGGA			
116251		CCTATTCTTG			
		ATCTTCCATC			
		TCCTGTTCCT			
		ACCTGCCATT			
		GATATCAGAA AGAAATTAAC			
		AAAGGCAGGA			
		AAAGGCATGT			
		GGACTGTGGG			
		TTTGCAGAAA			
		GTCAGTAAGT			
116801	CTAGATTGAT	AGGTAGACAA	GGGGTTAGAC	AGGTACATTT	ATATGTCACT
116851	GGAGAGCTCA	TTATATTGGT	ATAAAGTTAT	TGTGTCACAT	GTAAAGTATG
116901	ACATGGGGGA	ATTGGGGAGG	AAGGAGTGGA	ATAATACTGT	CGCTGCTAAG
116951	ATAGGCATTG	TGATATGGTG	CTTAAACCTG	CAAGTAAAGG	AAAAGAGTAT
		TGTCTTTTTC			
		GGGTTGCTGG			
		GAATTAACTA			
		GCTAGAGGAT GAGAAAACAT			
		ACCACGGACA			
		CGAACACTGA			
		CCATCTTCAT			
117401	ATATAAGGCT	GTCTGCTTGG	TAATTTAAAC	CCTTGGCTTA	TAGTCTTTTC
117451	AGTGAATTTC	TTTCCTTGCA	<b>AACTCGAGAG</b>	TTGGAGTCTC	ACGACTGCCC
117501	TTGCTTCACC	AATTCCCCAG	CTAGAGACAA	AAGACCTTCT	TGGCCTCTGA
		CCTTGAGATT			
		GGAATGAAAG			
		ATGTGGACTT			
117751		TGCCAAGAGC TTTCACTCTA			
		GGACTCATCA			
		TGAGAAGGGA			
		ATTTCCTTTG			
		GTGGTTGAAC			
118001		CTTGCCCTTT			
118051	TCTTGAGAAC	TTCACTTGAT	GCTATTTCTC	AGGAGATGTT	TAGGTCAGGT
118101	TGTCCACCCA	GGTATAAAAG	AGAAAGAGGA	ACGCTTATCC	CAGTCTGCAA
		CATGGTCTGG			
		TATATATAAA			
		CATGCCTAAT			
		AAGCGAGTCT			
		TATGGAAGAA TGGGTGAACA			
		AAGGTCAGGC			
		GGAGGCAGAA			
		GAAGTTGAAA			
		ATTTGTAACA			
		GTTAGGAAAC			
		GATGTATTAT			
		ACTCACTACT			
		TTGTATTATT			
		TGAGGGACTC			
		GGTCACTTAT			
		TCCATCTCTA			
		TACATGCCTG TGAGCCCAGG			
		CCAGCCTGGG			
		AGATTACAAC			
		CGCCCAAGGC			
119251	GAGACAGGGC	ATCTTTCATT	CCTTTGAAGA	ACCAGACTCC	TCATTGGTTC
		AACCTCATGG			

FIGURE 3, page 31 of 57

119351	GGTGGACAAA	CTGATCAAGA	AGACAAACCT	GGCCTTGGTT	GTGGGGACCC
119401	ATTCCTGGAG	GGACCAGTTC	ATGGAGGCCA	TCACCGTCAG	TGCAGGTGAG
119451	AAGTGTCTCA	GGCTGGCCTT	CCTCCCACAA	CCACCCAACC	TCTGAGAAGG
119501	AAGCGTAAAG	CCACGTTAAC	AGCCTCCCAC	TCCCTACCAA	CCCMMCMCMC
119551	TTCACTCTTC	CCAGCTCTGG	MCCCTGCCAG		
119601					AAGAATCATG
		AAAACATGGT			
119651		AACAATGTAC			
119701					CAGAGTCTTG
119751		AAACTCAATA			TTTCTTTTCA
119801	TGGGTTTGGT		TGTAAAATGT	GGGACAATTC	TGATTTAGAG
119851	ATGTGGGAGT	TAGGAGTTTA	TAAAATGTGT	TGCATTGACT	CTCCAACAAA
119901	ACACTCTGGA	TGATTCCATA	CCCCTCCCTC	GGCATTTACT	GACAGGCTCC
119951	CTCAGTAGTG	ACCCACAGCA	CAGCCGGGAG	TCCTAGCAGC	CTGAGGGGAC
120001	TGCTGGTTGG	AACAGGGACG	GAAAAGGTCT	CCCAACCACC	ATCACTATCA
120051	CCTCTCAGCA	CCACTGAGGC	CTCCTGGCCT	TGTCTTTTAT	TGAGAGACTT
120101	TGTTGTCATA	GCAACCCACA	GGGTCATATC	CCCAAGGCCC	CAGAGCCAGA
120151	GCAAAAAGAC	AGCCAGGAAG	AGAGGTTTGC	TECTECTECT	GCTGCTGCTA
120201	CCCCACTTTT	CTCATCACCT	CCTTTACATC	中で中でである。	CCCCCCCCACA
120251	GACCTGACTG	TGCCCCTCAA	CACAATAAAC		
120301	TACCCTGCTC				
120351			GGAGGGGACT	GAGGTTCAGA	AATAAGAGAT
120401		AGCTTACAGA			GAATGAGAAC
		TGACTCCTGT		CCCAGCTTCT	ACCGGTTATG
120451	CCAAAACATG	ACAGAAGTTG	CCGTTGGCAA	GGCACAGGCA	
120501		TCCAGGGCTG			ACATTTCCTG
120551	GCAAGGACAA		TCCTGCTTTT	TCCCATGAGA	TGTTTGGAGG
120601	AGGGCACTGG	CTCTGCAGTA	TATTCTCGTG	ATCTGGAATG	ACAGCCATCC
120651	CTCAGGGGAC	AGATAATGAC	CAGAACCACA	ATGGTTATTG	CAGCAGTCAG
120701	GTCAGAAAAT	TTGAGAGGAG	CCCTGCTGGC	ATCCAGTGAA	GAGTGGCCAC
120751	ACCGAACTGA		CTCCTTAGAC		AGCCTGTGCA
120801	TTCTCCTTTC			TTAAGTTCTG	
120851	GCAGAACATA	GAGTTTTGTT	ACATAGGTAT	ACACGTGCCA	TGGCGGTTTG
120901	CTGCACCCAT	CAACCCGTCA	TCTACATTAG	GTATTTCTCC	TAATGCTATC
120951	CCTCCCCTAT	CCCTCACCCC	TGACAGGCTC	CAGTGTGTGA	TGTTCCTCTC
121001	CCTGTGTCCA	TGTGTTCTCA	TTGTTCAACT	CCCACTTATC	AGTGAGAACA
121051	TGCAGTGTTT			GTTTGCTGAG	
121101	TGCATCCTCC	TTTCTTTCTG			
121151	CTTTCTTCTC	TTTCTTTCTG	CICCACIGIC	TIGICCCICI	TAATCTCCTT
121201	TGCCCTGCAT	TTCCTTATTC	CCIGGCCCIC	CCCCARAGE	
121251		TCAAATTGAC			TTTCCCCACT
121301	ATTTTCTGGC GGAGTCCCCT	ACGCTGGCCC		CAGCTGCCCA	
		TCTAGCGGAT			GGCTTGACTT
121351	TCTCATGAAT	GATTATCTGA			CTGTTTATCT
121401	TGCCTTCAGC	AGGGGATGAG			GGAGAGGCTG
121451	CCCTCCTGCT	TTGACTACGT			TCTGGAAGGT
121501	GCTGTTTGCC	TGTGTGCCCC		CTGCCACGGC	TGGGCCTGCT
121551	TCGCCGTCTC	CATCCTCATC	ATTGGCATGC	TCACCGCCAT	CATTGGGGAC
121601	CTGGCCTCGC	ACTTCGGCTG	CACCATTGGT	CTCAAAGATT	CAGTCACAGC
121651	TGTTGTTTTC	GTGGCATTTG			AGTGAGAGGT
121701	GCTTGAATTT	GCAAAGAGGA			CCCTGGACTC
121751	CATCTCATTA	TCTTCCACAC	CATCTCAGAT	CTGAACTTAA	
121801	GCCCTTAAAG	TGCACAAAAG	TCAATCAAAG	AGATGAATAA	TGACATTAGT
121851	AATGACAGCT	AATATTTCTT	GAGCACTTTC	AATGTGACAG	ACACCATGTG
121901	TGTTCAGCAA	TTTACACATT	TACATTTTCC	CCCTCTAATC	TOTOCCANANC
121951		TAGGGTAAGT	TATTATCCCC	ACTTCACACA	CARACARAG
122001	GAGGCCCACA	GAGGTTAAGC	TACATCCCCA	ACTICACAGA	CAAAGAAACI
122051	AACCTCCACA	TTATGTGAGT	ACACCACAAA	CACTCAAATT	CCAATTTCTT
122101	AGATATTGTT	CTCCTTCTAT	TTACCTCTCC	CAGIGAAAII	AAAAGAATGT
122151	ATTACCCACC	TCANACATAN	CARACCACAG	CGATCTCTGA	GAGGTTAAAG
122201	CCTCCTCTAC	TCAAAGATAT	CAAAGGAGAA	ATGCCCACAT	ACATTCTTGG
122201	CACCCAAACC	TTGGAAGGAC	ACTGTGAGTA	CAAAGTATCT	CCTAGCAGGA
122231	CAGCCAAAGG	AAGTTCCACA	GCTTTTATCT	TTTTATAGGA	TGAATTACAT
122301	ACTOTTOTT	TTTCTTAGGA	ACACTCAGAG	ACAAACAGAA	AGGAGCGGAC
122351	ATTCCTTTAC	TCATTGAACA	AATATTTACT	GAGCACCTAT	TATGCCTGTT
122401	ACAGTATTGT	GCTAGTTTTT	GGGACTATAG	TGAAAGGCAA	GATACACATG
122451	CTTCCTTCTC	CACGTGGAGT	TTATAATCTA	CTGAAGGAGG	CAACTCTCAA
122501	CTACTGTAAT	TAAAGTTATC	TTGTTAAATC	CTAGGAAGAA	AAAGAAAAGG
122551	TACTGCATAC	GGAAGGAAGT	TGGGCCTGAA	TGTAGGAGTT	AGCAGGTAGA
122601	CAGGGGCTGC	ACTAGCCCAG	GTTCTTTACT	TAATTCAGTT	AGGGGCTTTG
122651	GGGCCTCTGA	ACTCTGAACT	TCTGCCAGGG	AGCTGGCATC	CCAGTTGCCC
122701	CAGAAAGAAA	CAGAGCACAT	CCTCCTGCAG	GGAAGTTAGG	CTGAATCTCA
122751	TCAGACAGGA	CTTTTCTGGC	TGGGCCAAGG	GAAATCTTTC	CTGTACCAAC
122801	CAAACATATC	CTTCAAGAGA	GTAGCTCAAT	TCACATCANA	TTCTACCAAA
122851	ACCTCTTTCC	AAAACCCCAG	CGCAGGCCAC	CCCTDTTTTTT	TCTCCAMMAC
122901	TGATGCAAGA	GATTTAGCTA	TCGTGCAAAM	CCDTCXCXXC	TOTOCHI I AG
122951	AGATGGATGA	TCCCAGGAAG	CCCCCCAM	CACATCAGAAG	CMCAMCMCMC
123001	TTCTCCAAGC	CTTGGGGGAC	CTCAACTAT	ACACCCCCT	GREENTETETE
123051	GGGGGAAACC	DTACACCTCC	CARCARTE	AGAGGGGAGG	GAGGAAATAT
123101	DCDDCAMAGC	ATAGAGGTGG	TATAAAAAA	CAGAGGATCA	GAAGCAAAAA
122161	CCATACCCA	CAACAGAAAC	AAAAACAAAC	AAACAAACAA	AAAAACAAGG
*53131	CCMINGGCAA	GAAAGGGTAA	GAGGTTTTCT	CTGGGAGATC	TAAAAAAAAT
		~~~	~~		

FIGURE 3, page 32 of 57

	GGCAATAATG				
123251			GCTAACAACA		
123301	ACCCTGTGAT	CCACTCATCT	GATTTAGTGG	CTTTGGCTGA	AGCTCTTTGG
123351	ATATAGTTGA	AGGTACGGAA	<b>AGGGTCCTTA</b>	CATGAGGACT	TTAGGGTCAA
123401			GTGACCTTGG		TGACCCTTAT
123451			AAGAATTGGG		CTGACAGTCC
		TACAATCTGT	GCCAAGATCT		
123501			_		ACCCTGCAAG
123551			ACCACAAAGA		
123601	GTCCACAGTT		GTCCCCATCA		
123651	TGTCCCCAAA	ATCCAGCACC	TCACCCAGTG	CTCAATCAGT	AGGCATTGCT
123701	CAATAACTGT	TGGTGGTTCG	TGAATAAATG	CCCCATATGA	CAGTTAAAAT
123751			CTTCCCAGGG		
123801	ATATTATGGG			AAAAAGTAGT	
					CAAAGGAGAA
123851			TATCAGAACT		TCAGAATCTT
123901	TCAGATCACT	GCAGATGAGG	AATGGGAAGC	"CCAGACTAGG	GATGTGACCT
123951	ACCCAGGGCC	ACACGGCTTG	CTTGCGGCAG	AACTAGGAGT	TAGGAGTGGC
124001	CCCCTAGCCC	TTGTCTCTCA	TTCCTGGGTT	CAGCCCACCA	GCTCAAGCTG
124051			ACAAGCCCTG		CCTCCTACCA
124101	GTTCCCATGT		TTTTCCAGAT		
124151			CAGACGCCTC		GTGACGGGCA
124201	GCAACGCCGT			GCCTGGCCTG	
124251	GCCATCTACT	GGGCTCTGCA	GGGACAGGAG	TTCCACGTGT	CGGCCGGCAC
124301	ACTGGCCTTC	TCCGTCACCC	TCTTCACCAT	CTTTGCATTT	GTCTGCATCA
124351	GCGTGCTCTT		CGGCCGCACC		GCTTGGTGGC
			CACAACATGG		
124451			CACTAGAGGC		
	TCTAAGCCAC		CTCCAGCAGG		
124551			CCAGTGATCT		
124601	GAGAGGCAGC	ATCAGGACCT	AAGCCCCAGG	AACTTCACCC	AACTTAGGCC
124651	CTGGCAATTA				
124701			TTTAATTGAA		
	AAATCCACCT				
			TCCCCCTCAT		
124801			TCTATTCCAT		
124851	TCTCATTTCT	TTGGAAGCAG	GGTTTCTCCT	TETCTGCCCAA	TTCCATATGT
124901	CCCTATTATC	TCACTCAGCT	GACAAGACGT	GAAAATGAGT	CACATTCATG
124951	TGGCTGGGGT	GGGGTTCTTT	TTTCATTGTA	ATCATTATTG	TGGTTGCTTT
125001			CTTATTATTT		TTTTTTTCTG
125051			ACAAAGGAAT		
			CACATGCTCT		CTTGGCTCCA
125151	TCAAGATCCA		TCACTGTTTT		TTGGGAGGAG
125201	TGATGGTGTT	GGGGTAGAAA	TAAGCTCACT	CACCCACGCA	GGGTACTAAA
125251	GATCTTACAG	GAGCTTCAAC	TGGAGCAGGA	GGAGCTTTTT	ATGCTTATGT
125301	TGAATCAAGT	CAGATACAAA	AAGCAATTGT	CCCTCTTTGC	CCAAGCCTTT
125351	CCAATTCTGT	GTGTCTTGTT			ATCCTTCTGC
125401	AGGAAGACCC				
			AGAGATGGGA		
125451	GACGACAAAG		AGAACAAAAG		CTTCTTGATT
125501	ATCTTTGGCT	TTGTACCTGA	GGCAGGAGAG	AAGAGATGTC	CAACCAGTGA
125551	GATCTTTAAG	AGAAAAGTTT	GTATTTTAAA	TGTCAATGTG	CCTGAGAAAT
125601	GTCAGCTTCA	CCACGCTCTT	GCTTCCTAAT	GCTCTATACA	AAGAGGGCTG
125651	ACTATATTTC	TTGAAGTGGT	GTAAAAACTT	AGAGATTTTA	TAAGAGAACC
125701	AGGGGCTCCC	TTCACCTCTC	CTGGTCCCTC		TGAAAGCATT
125751			ATTCCTCATT		-
					TCCTGCTTGT
125801	TCTTAAACTT	CATGAAGCTA	TTTTTCCAGC	CTATGGGGTA	GTTCTTGCTC
125851	CAGTAAGAGG	AATCTTAGTT	GTCATAATCC	CTTGGAGCCT	GGGTTTTTGG
125901	AGAAAGAGAT	CTCCGTGCCC	TACAGACCTT	TTCTCAACGA	ATGTGGGAAG
125951	GACCTGGCTT	TAAAACACGC	ACACAAACAC	ACAAATAAAC	AGACATAAGA
	TGTCATCACG				
	ATACATTTTC				
	TTAAACACAT				
	GTCATAAGCT				
	AAAAAAATAA				
126251	TGTGTTTCTT	CAGTGAAAGG	TCCAGGGGGC	CACTGTGGGC	TTCTTGTGAG
	GAGACGTGAC				
	TGTGTCAGAC				
	TTGTCAAAAC				
	GCTGTGGTAA		AGGAGGTTTC	TGTATCAGAA	AGGCATTGGC
	CGTGACAGAC	TC			
(SEO I	D NO:3)			• •	

## FEATURES:

Start: 2010 Exon: 2010-3793 Intron: 3794-109509 Exon: 109510-109613 Intron: 109614-118338 Exon: 118339-118463 Intron: 118464-119345

FIGURE 3, page 33 of 57

Exon: 119346-119445
Intron: 119446-121409
Exon: 121410-121685
Intron: 121686-124128
Exon: 124129-124502
Stop: 124503

SNPs:

SNPS:						
DNA				Protein		
Position	Major	Minor	Domain	Position	Major	Minor
378	С	т	Bound OPE/EIN			
742	T	_	Beyond ORF(5') Beyond ORF(5')			
2005	Ċ	T	Beyond ORF(5')			
2381	A	ĉ	Exon	124	TT.	m
5165	c	T	Intron	124	T .	T
5402	A	Ğ	Intron			
6794	T	c	Intron			
9883	A	G	Intron			
10210	T	С	Intron			
12220	T	G	Intron			
13842	G	A	Intron			
14200	С	A	Intron			
15878	G	T	Intron			
16030	A	G	Intron			
16292	T	C	Intron			
16506	T	G	Intron			
17953	C	A	Intron			
23832 25001	C C	G	Intron			
25141	A	A G	Intron			
25191	A	G	Intron			
26147	-	AG	Intron Intron			
27400	A	G	Intron			
27401	A	T	Intron			
29278	c	T	Intron			
31437	A	G	Intron		•	
31857	A	G	Intron			
33155	G	A	Intron			
39487	G	С	Intron			
41449	T	С	Intron			
	T	С	Intron			
43256	G	C	Intron			
43967	T	C	Intron			
48604	_	A	Intron			
49560 52729	A G	T	Intron			
55031	A	T G	Intron			
55066	Ä	c	Intron Intron			
56912	A	Ğ	Intron			
58480	C	T	Intron			
61128	Ğ	Ā	Intron			
61320	G	A	Intron			
61444	A	С	Intron			
62641	T	С	Intron			
63023	A	G	Intron			
63051	T	C	Intron			
64989	T	G	Intron			
65929	C	A	Intron			
66694	C	G	Intron			
66755 66879	T T	A C	Intron			
69156	Ċ	T	Intron			
69280	č	T	Intron Intron			
70647	c	Ť	Intron			
71867	Ċ	T	Intron			
71900	Ċ	T	Intron			
71901	G	A	Intron			
72369	С	T	Intron			
72992	T	G	Intron			
73154	-	T	Intron .			
73164	_	T	Intron			
74149	T	A	Intron			
74171	G	A	Intron			
74918	A	G	Intron			

FIGURE 3, page 34 of 57

wo	02/33086	;	
	_		-
75386	G	A	Intron
77751 78264	G G	A T	Intron
80986	T	A	Intron Intron
83609	Ĉ	Ť	Intron
85271	Ğ	Ť	Intron
87770	c	T	Intron
87837	T	С	Intron
87866	С	T	Intron
88238	A	· C	Intron
89219	A	G	Intron
89331	T	C	Intron
90794 92404	A C	G	Intron
92672	A	T C	Intron Intron
92684	A	G	Intron
93132	G.	č	Intron
93537	A	T	Intron
93557	T	С	Intron
95067	С	T	Intron
96000	T	С	Intron
96877	G	T	Intron
97271	A	c	Intron
97470 97518	G G	T A	Intron
98476	C	Ť	Intron Intron
98779	c	Ť	Intron
99218	č	Ġ	Intron
100538	C	A	Intron
101045	A	С	Intron
101232	С	G	Intron
101266	G	A	Intron
101290	A	G	Intron
101326	G	A	Intron
102342 104489	C C	A T	Intron
105266	A	Ğ	Intron Intron
105338	T	č	Intron ·
105570	C	A	Intron
105928	. G	A	Intron
106459	G	С	Intron
107710	С	G	Intron
108062	G	A	Intron
108214 108364	G C	A A	Intron
108657	T	A	Intron Intron
109746	Ċ	Ť	Intron
111484	Ğ	Ť	Intron
112879	A	G	Intron
113245	С	T	Intron
113265	T	С	Intron
113497	С	G	Intron
114486 114686	G	T	Intron
114885	Ť C	C A	Intron Intron
115600	G	T	Intron
115668	A	Ĉ	Intron
115745	A	Ğ	Intron
117230	A	С	Intron
118908	A	G	Intron
120430	С	A	Intron
120830	A	T ·	Intron
121926 122102	T	C	Intron
122102	G T	C C	Intron
123366	Ċ	T	Intron Intron
124947	c	Ť	Beyond ORF(3')
125010	A	Ğ	Beyond ORF(3')
126043	T	c	Beyond ORF(3')
126064	-	G	Beyond ORF(3')
126283	С	G	Beyond ORF(3')

Context:

DNA Position

FIGURE 3, page 35 of 57

PCT/US01/32152

378 TGGCATGTACAAAGGTCCTGGGGTGGACAGTCACTTGGTATAATCCAAGAGTGAACCTGA AGGCTATTGTTGAAATGTAATAAGGGAGAGAGTGACGGGATGAAGGGGGATGAGTGG GAAGCAGTGAATTCCTGCAAGGCTTTGAAGGTCATGGGAAAGAATTTGGTCTTTATATCA AGAGCAAGAGAGACTACTAAAGGGCTTCAAACAGGGGAGCGATATGCTTAAGTCTGTTT GTTTGTTTTTTAAAAAAAGATTACGGTGGCTATATGAGGAAAGTGGAATTGAGAACTAG (C.T) GAGACTTGGAGTGGGGGCTCCATTAGGAGGCTACTGAAGTAGATTCATGAGGTAAGGAG TGATGCTGGCCTGGGCTGGGATGATGGTGGTAGAAATGGAGAAAGAGTTGATAGGATTTA GTGATTGGATAAGGGACAGAAGAGAGATGAAGGCTTTCAGACTAACATCTGCTTTCTAAC ATGAGTAACTGGGTGGCTGAAGATGCTATTTTCTGAGCTGGGAAACAGGAGAAAAAGGAG CAAATATGGGGGATGAAGACTTTGAGTCTTTAAGGTGCTGTACAAACACAAATCAGCATT TGGTGGCCTGGGCTGGATGATGGTGGTAGAAATGGAGAAAGAGTTGATAGGATTTAGTG 742 ATTGGATAAGGGACAGAAGAGAGAGATGAAGGCTTTCAGACTAACATCTGCTTTCTAACATG AGTAACTGGGTGGCTGAAGATGCTATTTTCTGAGCTGGGAAACAGGAGAAAAAGGAGCAA ATATGGGGGATGAAGACTTTGAGTCTTTAAGGTGCTGTACAAACACAAATCAGCATTCCT TGAGTGGTAAGATGAGTATAATAGTTTCAATTGCATTTCATCCCATTCTTCTGAGCTC AAGCTCACCTTTTACTGGTTTGAGGCCAGTAGATGAAGCTGCATATCACCCCCAAAATCT TGTCTCTAGTTTAACAAACTTATTTGAGAGACATTTGCATGTTTTATTAATAATGATTT 2005 TTTCCATCCCCACTATTCCCACCTATTTCAAGCCATTTTCAACGGAGTCTCCACCAGAT GGTTTGGACKIACAGAGCAGCTATTTTTGCCTCCCATTGACATCTATTTTTCCAAGTGAGA GACTGCCCCATATUTTAUTIAL AATATGTCACTGGAGGTGAAGCATCAGTTGTATTGGTGG GAACCTGCCCTTT / TOTAL TITTCCTCATGCCTTTTCCTGCCTCTCTGATCTTTTC TAGGTCTCTCAUCTAT A DOA DUACAACTGGTGCTGCAATAGAAGCCAGTGGCTAAGTCT [C,T] GTGTATGGC&TSUTTAASGTTGCAWCCTCTCACCTCTGCCTTCCTCCATTTTGGGCTGGT TACCTTTGTGCTCTTTTTGAATC+TTCTTCGAGCAGAGGCTGGTGGCTCAGGGGACGTGCC AAGCACAGGGCAGAATAATGAGTICTGTTCAGGGTCATCGGACTGCAAGGAGGGTGTCAT CCTGCCAATCTWITA 12000A TAACCCTTCCCTTGGGGACAAGATTGCCAGGGTCATTGT CTATTTTGT GCCCCTGATATACATCTTCCTTGGGGTGTCCATCATTGCTGACCGCTTCAT 2381 CCTGAATGGTCTTCGAGCAGAG ACTGGTGGCTCAGGGGACGTGCCAAGCACAGGGCAGAA CAATGAGTCCTUTTUAUSGTCATCGGACTGCAAGGAGGGTGTCATCCTGCCAATCTGGTA CCCGGAGAACCCTTCCCTTGGGGACAAGATTGCCAGGGTCATTGTCTATTTTGTGGCCCT GATATACATCTTCCTT XXXXTGTCCCATCATTGCTGACCGCTTCATGGCATCTATTGAAGT CATCACCTCTCAAGAGAGGGACGTGACAATTAAGAAACCCAATGGAGAAACCAGCACAAC [A.C] ACTATTCGGGTCT(X)AATGAAACTGTCTCCAACCTGACCCTTATGGCCCTGGGTTCCTCT GCTCCTGAGATACTCCTCTTTAATTGAGGTGTGTGGTCATGGGTTCATTGCTGGTGAT CTGGGACCTTCTACCATTCTAGGGAGTGCAGCCTTCAACATGTTCATCATCATTGGCATC TGTGTCTACGTGATCCCAGACGGAGAGACTCGCAAGATCAAGCATCTACGAGTCTTCTTC ATCACCGCTGCTTGGAGTATCTTTGCCTACATCTGGCTCTATATGATTCTGGCAGTCTTC 5165 TTCCTCTGAATGACTGAACATATCCACAAATAATAAGCGTGGCAGGAGATGGTGTGAAGA GTAAAAGGAGCATATAGGAAGTTGTGTGTGTGGGGGTGTCTGTTTCAAGAACCTGCTAATT GGAAAAGTGGGGASCCATAGAAGCTAGGGAGAGGTGTCCTAGGAGTGCTTCTGCCCAGGT CCAGCCATGAGACAGAGCTCAAAAAGAGCTGGGCACTGCTGGTGACAGAACTGAGTGACC [C,T] GGGGGATCCTGCATCTTACTCAATCCCTTCTTAATAATGTGACTTGGGGCAGGTC ATTTATTGGTTCTGGAACTTAACTTTCTGATATGCAAACTGGGAATAACAATACTTTCCT TGCCTGGAGGCAAGGTCAGTCCTTTTTGCAGTTCCTTCCAGCTCTAAGATTTTCTGAACC ATAGACATAAGCACTCAGTGTAGGTCATATTCGCACTTGCCAAAAATGGATCAGGGAATA TTGTCTCCTGAAGGGAAATGGCCATTGACAAATTGATTATTAGAGCTCTGTTTAGTCAT 5402 GGTCCAGCCATGAGACAGAGCTCAAAAAGAGCTGGGCACTGCTGGTGACAGAACTGAGTG ACCCGGGGGATCCTGCATCTGTTCTTACTCAATCCCTTCTTAATAATGTGACTTGGGGCA GGTCATTTATTGGTTCTGGAACTTAACTTTCTGATATGCAAACTGGGAATAACAATACTT **AACCATAGACATAAGCACTCAGTGTAGGTCATATTCGCACTTGCCAAAAATGGATCAGGG** [A,G] ATATTGTCTCCTGAAGGGAAATGGCCATTGACAAATTGATTTATTAGAGCTCTGTTTAGT CATTTTGCTGGGAAGGATAATCATTTGTTAACGTAAGTAGAAACCTGTGCCTTCTGGAGA CTTCACTGGGAAAACAAACTCCATGGAATTTCACATGATTATCGCGATGTCAGTGTGGAA GAAGATATGGTAAGGCATTAAATGACATTAAGACCACAAAATTTGCCATAATTTGACGGA 6794 CTCATAAAATATTAGAGCTAGAAAGGACCTTAGAATATCTTCTGCAGTCATGGTTCTTAA ATTTTAATGTGTTGCTCAATCATCCAGGGATCTCACTGAAGGGCAGATTAGGATCCAGGA

FIGURE 3, page 36 of 57

GGTCTAGGGGAGGGATTGAGATTCCGCATTTCTAACAAGTTCTGGATGCTGCGGGCCCCA ACTTAGAGGTGAAAGGTTCTGAAGCTCTTGACCAAACCAGGAGACCCAGCAAAGAAGTGG

TTTTTCAGACAACTTGCTTAATTGAATAATGATTGTTTGCTCTTTAATTCCAACTTTCAA [T,C]

10210 CAGATGCTCAAGTACCTGGTATAAAATGGCACAGTATTTGGCATATGACCTAGGCATATT
CTCTCCCATATACTTTATTTATTTATTTTCGGGACAGAATCTCATTCTGTCGCCCAG
GCTGTCACTCGCTTATTGCAACCTCTGCCTCCCAGGGTTCAAGCAATTCTCCTGCCTCAGC
CTCCTAAGTAGCTGGGACTACAGACGCATGTCACCACGCCTGGCTACTTTTTGTATTTTT
AGTAGAGACAGAGTTTCACCATGTTGGCCAGGCTGGTCTCAAACACCTGACCTCAAGTGA
[1, C]

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12220 ACATCCAAATAGTAACTTAATATTCCAAATATGGCTGCAAAACAAATTGTCGATTATGGA
TGACTACTACTGCCATCTCTCCATACCAGTCCATCTTCTGCCAGGCTGTTTGGTCTTGAT
TTGTCGACCTTTTAGGTTTCTCCCCATGTATTCCACATGACCTTCACCAACCCCACTTCT
ATCTCCAAACGTCTTTCTGAGTTGTGGGGATGCAGATGTATTCTGCCACCATCACAAGGG
CTAACCGAGCCCTGGCTGCGGATCTTCATTGTTGTTCACATTATTTCCATTCTTACACCC
[T,G]

ACTTCATGTTTGTACACTATTTTCTTACATTTGCTGCTCTCTTCTAAACATTCTTTGCTGC
ATCCACTTTTTCTCTATTTGTGCTCTAGGTGCTGCAGAGGCTAATGCTGGGTTTCCTTTC
ATTCCTCCTTGCACTCAGCACCTCCCTTCTCAATTCCTTTTGCCATGTCTCCACTTTAAA
TCTTAACCTACTCCAGATAGTCTTTTCCTTCACACTATTGGCATCTGTGCTTGGGTTGCT
TTCAGTCTATTCTCTGATCTATGATTTCTTTGCATGATCAAGAAGGTGCCATGAAAGGAT

13842 TCACTTTCAAAGCCTCTTTCTGGGTTTGGATTTCCAGAGCAGCCTGTGCTGTAAAGCAAG
ACAGAAAGCTTCCCTGCCATTCATGCCTGCCAGGGATAGAATGACAGTACTCCTGAGGCT
CTCCCTCCCACCCCTCCCTGCTGGACAGCTGATCTGCTGGACTCAGCCAGAGCCAGCA
GCCCCCCTCTTTATCCTAGGAGCTGCAAACTTGATGCCTTTCCAGGAAATCCCCAGAA
GCTGGAGTATCCTCATCTACATGTGGCACAGTGTATGGTTGTGCAGGTGCTCATGTCCC
[G\_A]

TTGCATAGGACTGGGGTGGAAAATAGGGACCGTCCTTTTGTGTCAGCTCCAGTCAATGAG
TAGTGGCCATCCAGGGGCCATCTTGGAAAGGACTTGTGAGGCTGTATCTGCGCTCAGTT
GTAGATGTGAGAAAAAGGCCAAATATCTGCCAATCCTAGTCCTGGGATTCAAGATAGA
AAGAACTGCATGGAGTGAAGAAACTAGGAGTCTCCATTTCACTGAGATGCATAAGAATGA
AATTATTGTCACTATTTCTTCAATACTGGGCCAATCCTAATAAGAAAACCCTTTTTGAGT

14200 GAGTAGTGGCCATCCAGGGGGCCATCTTGGAAAGGACTTGTGAGGCTGTATCTGCGCTCA
GTTGTAGATGTGAGAAAAAGGCCAAATATCTGCCAATCCTAGTCCTGGGATTCAAGAT
AGAAAGAACTGCATGGAGTGAAGAAACTAGGAGTCTCCATTTCACTGAGATGCATAAGAA
TGAAATTATTGTCACTATTTCTTCAATACTGGGCCAATCCTAATAAGAAAACCCTTTTTG
AGTCTCTTTTTCTTTATCCTACATATAACACAGAAGCTTTTTCTATTCCTGGATGAAC

15878 TGTGTCAAAATATCACTCTGTATCCATACATATGTATAATTATTATGTGTCAACTAAAAA
TAAAAGGAAAAAAATCATTTCAGTGTATTTACAAAACATATGTAACCATTAAGAATAATG
TTTTAAATTATATCTAAGGGTGTGATAAAATTACAGTATAAGATTGTGCTTGAAAAAGTG
CAATAAGAAGTAAATATGTACAGATGAGAAAAAGTGCAAAGAACTAAGTCCTAAGCAGAC
TATACCTTTCCTACTGCATGGTACTTCTCTGGCCTTTTGAAAAGATTTTGCACCCA
[G,T]

FIGURE 3, page 37 of 57

ACAGTATAAGATTGTGCTTGAAAAAGTGCAATAAGAAGTAAATATGTACAGATGAGAAAA
AGTGCAAAGAACTAAGTCCTAAGCAGACTATACCTTTCCTACTGCATGGTACTTCTCTGG
CCTTTTGCTTTGAAAGATTTTGCACCCAGCATGGCAAGTGGTTAGCAGAGGCAGCCATTC
TCACTTGTGCGTTGGCTTTGGGAGCCATATATGTTGTTCAGCTGGTGTGGAGTGGAAAG
GCTGCATGTTGTATTAATGCATTGTTAAGAACCTCTAAGAGTGATTTCTTTTGGGAAGTG
[A, G]

GACTGACGGTCCGAATGGTGGAAAGACAACTTTTAATCTTTTACTTTACACTTTGTGCAC
TTTTAAATGTTTAACATGAGCATGCATTTCTTTAATAATAAAAAATACAAAAAAATTTTAG
CCCTAGATCTTCTGATTTTAAACTGCATATTCTTTCTATTGTGTTACATATTTTAGCATG
AGAATAAGGTTATGAAGCTGGAAGTAGCAGGCTCCCTTTTCCTCATATGTAGGAAGTTAA
GAATGCATTCTACGTTTCTTCTTTAAGGAGTTGCTTCTTTCCTTTTAACATAGGGGTAA

16292 TGTTAAGAACCTCTAAGAGTGATTTCTTTTGGGAAGTGAGACTGACGGTCCGAATGGTGG
AAAGACAACTTTTAATCTTTACACTTTGTGCACTTTTAAATGTTTAACATGAGC
ATGCATTTCTTTAATAAAAATACAAAAAAATTTTAGCCCTAGATCTTCTGATTTTAA
ACTGCATATTCTTTCTATTGTGTTACATATTTTAGCATGAGAATAAGGTTATGAAGCTGG
AAGTAGCAGGCTCCCTTTTCCTCATATGTAGGAAGTTAAGAATGCATCTACGTTTCTTC
[T,C]

TAATATATATATAATACCAGGCAGGGTTATTTTTTCCTCAAGTCATTTTCTAATTTT
TTTTAAATGAATAGATAGAAGAGCTGAAGTAAGGGTCAGGAGCAAGAGCTCTGCTTCCTT
TTCCCTTGCTGGGCTTCGTTAGAGAGCCCATCATCTCCTCAATATGTCTCCCCAACTCTTCT
AGGCATTGGATGAGTTTGCTGCAGATACGAAACCCAACTTTGCCAGTCACTTCATACTAA
CAGGTGAAATGTAGTGGAGGAGCCTTTTGAAGACAGGGACTCAGCCCCCATTAGCCTCA

> TGTGGGTGCAGGCACCAACCAAACCCAGTTGGCACCGTTGTCTTTTCTCTGCAATGAT GTATTGAATTTAATAATGGAGGTATATGAAATTCAGAGTGATTGGAACTGAAGGTTTAGG GGCTTTGTGTAAAATTGATATGTAAGGGATTTGGAAGTAGGTGAGGGATTCTTCCCCAAT ACTTATTCAATTTTGGAGTCAAATAACCAAGCATTTACAAATAGCCAAAAAAGAAATTGA AAGAGGGTTTAATCCAATAAATTTTCATGCCTCATATGAACCACATCTTATAATAAGAAT

25141 CCCCTGTATAATCAGTCAGCCAAATGGAGCAGGACCCTGTGTTTTGTAGCTGATACAACA
GGGCAGCATCTCTAGTGAGGGGGCCAGGGCTTCTATTTCCTTCATTAAAAAATGAAACAG

FIGURE 3, page 38 of 57

CAGACCTGATTCCATATTTAGAGATTACACTTAGTTGCCACTGTGGGTGTGCAGGCACCA ACCAAACCCAGTTGGCACCGTTGTCTTTTCTCTGCAATGATGTATTGAATTTAATAATGG AGGTATATGAAATTCAGAGTGATTGGAACTGAAGGTTTAGGGGCTTTGTGTAAAATTGAT [A,G]

25191 TGATACAACAGGGCAGCATCTCTAGTGAGGGGGCCCAGGGCTTCTATTTCCTTCATTAAAA
AATGAAACAGCAGACCTGATTCCATATTTAGAGATTACACTTAGTTGCCACTGTGGGTGT
GCAGGCACCAAACCCAGTTGGCACCGTTGTCTTTTCTCTGCAATGATGTATTGAAT
TTAATAATGGAGGTATATGAAATTCAGAGTGATTGGAACTGAAGGTTTAGGGGCTTTGTG
TAAAATTGATATGTAAGGGATTTGGAAGTGAGGGGATTCTTCCCCAATACTTATTCA
[A.G]

TTTTGGAGTCAAATAACCAAGCATTTACAAATAGCCAAAAAAGAAATTGAAAGAGGTTT
AATCCAATAAATTTTCATGCCTCATATGAACCACATCTTATAATAAGAATTATGCTTTTT
CATTTCATACTCAGTTAACAAATATGATTTGTGAGCACCTGGTAAGTTCAGGGCACTAGG
CTGAAAGGGGTTACCAAATGTCTTCATTTAACAAAGTCCAGCTGAGCTCTTACAGGTACC
AGAACTGTGCCTGGGCTGTCATATGAAGATGAATGTAAGAGTGTGTCAGGCCTTCAAGAG

> TTTGTTTTACTTCCCATAAAAACTCTTTGTGTCACATGGAGGTGAATGGAAAGAGAGGCT GTGGCAACAGACGGGAGACTTTTCTGATATCAGAACCCAGTCCCATAGACCAGAATGTAT GCTTTCAATCCACGTTGTCTGGGTCCATCCTATTGAGTGCCCTGCCCCCACAGCGGGGTA TGGAGAAGAGTCAGACACAGCCCCAGTCCTCACGTAGCTCACAATCCAGTGGAGGAGACG GACTCAGAAACAGATAGAGATGAAGCCATGAGATCAGTACTGTCCGAGGCCATGGCCACG

27400 TAAACTTTACAAATCCTTAATTTGTAAAATGTGGGCAATGATAGTACCTCCTCACAGGAT
TATTACGAGGTTTACACGGAATACTCTCAGCTCATAATAAGCACTTGCACAGGCCTCATG
GGCTAGGCCCTCAAAACTTAACGCATCTACAGGCAACAGCCATATGAAAGGAATTTTATA
CCACCAAGTCAAAAAATCTGTGAGCACTGCTCAGAAGCAAAAAGCCTGTCTCAACAGGG
TCATTTAAGGGGTGGGCGAGCTACAGAGAAAAATGAGCCCCCACAGGGTAAGCTGGGG
[A, G]

27401 AAACTTTACAAATCCTTAATTTGTAAAATGTGGGCAATGATAGTACCTCCTCACAGGATT
ATTACGAGGTTTACACGGAATACTCTCAGCTCATAATAAGCACTTGCACAGGCCTCATGG
GCTAGGCCCTCAAAACTTAACGCATCTACAGGCAACAGCCATATGAAAGGAATTTTATAC
CACCAAGTCAAAAAATCTGTGAGCACTGCTCAGAAGCAAAAGCCTGTCTCCAACAGCGCT
CATTTAAGGGGTGGGCGAGCTACAGAGAGAAAGACTGACCCCCACAGGGTAAGCTGGGA

29278 ATACACTTCAGCAAGTCACCTAACCTGCAAATTTCAAGCATGTGAATCTTGGATCTTTCA
TGTGCTAGCTGTGAGACTTTGAGAAATGTATTTAATGTCTCTTTGCTTTCTACCC
ACACAATGGGTATAATAATGTCTACCATATATCTTTGCAGCAAGGTCTAAATGGGTGAT
ACATGCTGAATACATTTCCAACAGAGTCTGTGCAATGATAAGCTCTTTCCAAATGTTAGT
TAAAGCTAACCAACTAACCCACCAACAAACCAACCTCTTAGCCAGGACTGATGGAAGGAG
[C,T]

AAGTTCCATCCAACATCCCATTAAATATGTAATGTGTATTAGCACAGCGCCTGGCACTGG

FIGURE 3, page 39 of 57

GCAGGTATTTTCTAAGTGATAGCCAATGCGAAGCCTACTTTATTATTTTCCTCTTTGCTT AACCTACAAGGTGTCTAAGACCATTTGTTTGTCCACACATAGTAAGATAAACAGCACTGA GACTGTGGTCCTTTCTGCCCTGTGTCCTTATCCCACCTGGGAATCTGGAAAGCCAAGCCT AGACACACTCGTTCCACAAATGTTTACTGAAGCTTGTTCTATTCAAAGCACTGTACAGCT

TAACCTACAAGGTGTCTAAGACCATTTGTTTGTCCACACATAGTAAGATAAACAGCACTG
AGACTGTGGTCCTTTCTGCCCTGTGTCCTTATCCCACCTGGGAATCTGGAAAGCCAAGGC
TAGACACACCTCGTTCCACAAATGTTTACTGAAGCTTGTTCTATTCAAAGCACTGTACAGC
TACAAAGACCATCTTTTCTGAACTCCAAACCAGGCCACATGGTTGGAATAACTTCAAGTA
TGGAGACCAAGAGAAAAGGTGGTTGTTGTCAGCAAAGCTCTGAGTCCACACCTTCCAGGA
[A, G]

33155 ACAGTGCTGAATTTCACAAATTGCGAATTAGGAAATTGTTGCTCATTTTACAATTTGGT
TTCCCTCAGGATTCCTTTTAAGTAGCCAGCTACCCCAGTACTTTTGAAATATGACTTGCT
TATAAAAATTTGATAGGCTTGGCACGGTGGCTCACACCTGTAATCCCAGCACTTTGGGAG
GCCGATGTGGGGTGGATCACGAGGTCAAGACCAACATGGTGAAACCCTGTCC
CTACTAAAAAATACAAAAACTAGCCAGGCATGGTGGCACATGCCTGTAATTCCAGCTGCTC
[G, A]

GGAGGCCAGGCAGCTAGGCAGGAGAATCACTTGAACCCAGGAGATGGAGGTTGCAGTGAG CCAAGATCATGCCACTGCACTCCATCCTGGGTGACAGAGCAAGACTTCATCTCAAAAAAA AAAAAAAGATATATAAACAAGTTTTTATAATATTCTCAATATGAACTAGTAGAAAAAAAG CATGTGTTTTTAGGTCTTAGAGGCCTGGTTCCCAGTTTTATCTCTGACTCTAATGAGGTA TAGTATTACCTACATTGATTAGCCCTTCTATACTTCATAGGAGATGCTCCAAGACTGCTA

39487 CACTTTGCTCCATCCCTTGGCCTTCTGCAGTCCAAGCTCCATCTGAGATCATCCAAGGC
TTCTCTTCTGTGTTGATCCTTGGCCTTCTTGGAGTCTCTTCTCCCATGTTCTCCCACAAC
AGAGCATTCTCCTGACTGTTTTCATTCTGCATCTCATCATCATCATCTTTTTCTC
TACCATGCCCCATAAATTTGGGTGCTCCTGAGGGTCCTGTCCTTGTCCCTGCTTTCTTG
TTGTACAACCTCCTTGATCTACTTCATCTACTCAAGTTTGGTCCACAATTTCTATATTGT
[G,C]

AAGATTCAAATCTGCATCTCTAGCCATATATCCATTTGCCTGCTAGGCATTTCTACCTGA ATATTTTATAGGCATGCCAGTGGCTCTTACTCTATGGCTCTTACTCTAAGTCTAGACTAC AGCAGAAAGCAATGCTCTTTTTATTAAGGCATAGTGCCTCTTTCAGAATAATTTACAGCA TACAACCAGGCCTGCTGTGCAGCATTACAATTTGTCATTAAAACTCCATTCCTCTTGCCA GAGTAAATGAGCCATTTACAGCCAGGGCGCCAAGATGGACTGTTGTTATTTTTTCTGCCT

TCAGATTCCAGGACACCAAGTTTTCTGTGGGAGCTTCCCTAGGAATATAACTAAGGAATT
TAAATCAGGTTCAGCTCATGCTGTTACACTCTCTCCCACTCAGGCATTGGGTGTGGC
TTTTCCAAGCTTGAGAAGGGTGTGATCTGAGATGGCTTTGGGTATAGAGGGAATTATAT
TTAGGTCTACCCTGTATAGGAAAAAGTGCCTTCCCAAAGTCTCCCTGGCCTAAAGTATAA
GAGATATGTGTTGGGATTTAGACCCAGAGCCCAAGCCAATAATGGGACCCCCTTCTCACA
[T,C]

GTGGCTACCTCCTGCTATCACCACAACAGCTATCATACCCATAACTACAACAGAGGCCAA TTAACGTGGTGATAATTGACAAATGTCAAGACATCCTACATTGAGGCACACTGTGCGTTT TGCGTGAGCTTTTAAATTGGTAGGGAAGGAAAACTTTTATACCTACACCTATCATGGAAG GCAGAAGGTAAGAGCTAAAATAAAGGTATGCCAAGAACAAAGGCAGGAAAGAAGGGTTTT AACAACTTGAGGCCTGATCCATTGATTAGTGAAGAGGAAACATGTTCAAAAACCACTCTA

43256 AGAAAAACATTAGAATGGAGAGCTAACTCTTTGGAAATGGTCAAAGAACACGGGTCTAC
AAAACCGTCAATAAAGCGCTAAGATGCCTGGGCGGGGTCAAAAAAGTCTACCTGGGCGGGG
TCAAAAAGTCTACCTGCTCAGCATATGGGCCCAGACATCTGACCTTTACCAACTCCACA
ATAACCACTTCATCTATGGATCCAGTCTTGGTATCACCTAGTCGTTTTCAAGTAACA
GAATATTTGGTTCTCAATGGTAGGTGACTGGAATACAGCTTACTTTCTCCCACCCCTACC
[G,C]

CCAATCCTTTCTGCCCCCTTATAGTTTAATTTGCTTGTAAATTACTTGGGAATACATTTG
GGAGCCATTATAGGGAAATAGAAGGCAGACATGATGAACAGAATGCAGGGTGTTTTTTAT
TACTTCACATTGTGCTCAACAATTAGGAGGAATTCTAGAAGCCCCTCCCAGTGGCCAGGA
ATTGGTCATAGCATGAATAAACTCAATATAGGTTGAGTATTCCTTACCCAAAATGCTTGA
TACCAGAAGTGTTTTTGGATTTTTGGATTTTTTTTTGAATATTTGCATTATATACTTACC

FIGURE 3, page 40 of 57

> AAAAATGTAAAATTTTTGAAAAGAAAGCCTCATTGAAAAGAAATCCCTCTCCCCAGCTGG GCTCCCAGGCAGCCTCCTGCAGAACATCCTTAGCATTGCAGAGGTTGTTCCCATGGCAACC GAGTAAGGGGCTTTTTGTTTTCCTTAGAAGATTGAATCCTTTCAACCAGAAGGTAACCAC TGGTTCTTCCCCACAATCCACACTCCAAACCCCCTACCCTTATTTGACTACATGACTAGT TTTGCATTTATGGATTTTTTTATGCCTAATTGAAAAAGGCTAAATATACAGAAACTGAGG

49560 TGAGGGGTTATGAGACCATAGGCTCATTTTGGGGGGGGGTCTAAAATGCAGTATTTTTTGA
ACTGATATGGGGAAAAAAGACATTTCTGAATTGTTGCAGATTCTGGGCCGT
TCCAGCATAAGCACCTTTCTTAGAGTACTTGGGTTTGTGAAGTAGTCCTTATCCCCTCCT
TCCACTATTTTACATCAAGTTAAAATAGAGGAAGATGCCTAGAAATGGCCGTATAGACAG
AGAAAACTGCACTAAAACTCCCTCCGTCATGCCTGACTCCTCTCTAGACTATGACCATCG
[A. T]

GGGGCCAGAAATCATATCTTAAAGATCACTGTGCCTCCAGTACCCAGCACGGTGTTTAAT
AAATGTTTGTTGAATGAACGAACTAGTAAAATTTTCAAATCATTAGAGCTGAAGTATCCT
TTAAGATTCTTTAGTCCCTCATTTTACAGATAAGGAAGCTAAGGCTCAAGACATTGTGTG
GCTTGGCCAAAGGCACACAGCAAGCTAAAGGCAGAGGAGGACAGGACCCGGCTGTCTCA
ACCCCCTGGCTGCTACACTTCCTGCAGCATTTCTAATTCTTTTACCATTCTTGCGAGGGA

52729 CCAATGGGGAAGCACCAGGGTCAGCCGCAAGGCAGAGGAAGCAAGAGGAAAACATGGACA
AGAGGCTCTACTGTGGATTCAGTGGCAAAGAATGGAGGAGGAGTAAGCAGGTTTAGG
ATTATCGGGTTTGAATGACTTGATTGAGCTGTAGGGTGTAGAGACTGCCTCTACTGTCTG
GCACCAGGGGTAATTAGGGCAGCTGGATAGTGGTCTGGAGTGTAGAGAGCTCCCTAAAGGA
GGTGGTTGGAGGTGTAGGTTTTGGATTGGTTGATCTGTATATGAAAGGTGCACGTGCAGG
[G. T]

55066 CACTTTCAGGCTCCCTTAGGACAGCCTCCACCTGCTCCTACTGTGCTTCCCATCGTCCC
TCTCCTCAGGCACAGGCTGAGGAGTAATAAGAGCACCTGATATGTGTCAGGCCTTACTGT
GTGCTAGGAATTGTGCTAAGTACTTCCTATGAATTTTCCTTATAATAACTT
TGTAAAGTTAGAGCCATTATTCCAGAAGGGAAAACCGAGGCAATGGGAGTCAAAGCAAAG
AATTTGGGCTTTTAACCATTACACTATTTTGCACAAGTAGCCAGTAATGAAAAGGCTGCT
[A, C]

TGAAATACTTTAAAACTTGTAGCTTCCTTCAGCACAGAAGTGGCTCTCTGAACCAATTTT
AAGCAATCCTGGCTCTATCTGTGCATGTTGATTAGCCTGTGGTTATAGTGTTAACAATT
TAGTGATTCACCTCATTTTTAATCTCTCTTTCCCTTTAGCAGGATCATTTTCTCTGTGTT
AAGGGATCAACATTGAGGTAAGAATGGCTAAATAATAGCATCTTCTGGAATACAAATGAC
TTTATAAATAAAGAAGATAAAAGGAAGAAGTAGATTTCTCAGCTCTAATACACTT

FIGURE 3, page 41 of 57

[A,G]

GCAAATGCCATATGCTTTCTCCTGCGTGTACTGGTCAGGCCAGTTCTAGATACAATCATG CGCTGCATAATGATGTTTTGGTCAACAGTGGATTGCATATGTGACGGTAGTCCTTTAAGA TTATAATACCATATTTTTGCTGTGCCTTTTCTAGGTCTAGATATCTTTAGATACACACAT ACTTACCATTGTGTTCCAATTGCCTACAGTTTCCAGTACAGTAACCTGTTGTACAGGTTT GTAACCTAGGAGCAATAGGCTATACCATACAGCCTAGGTGTGTAGTAGGCTATACCACTT

58480 ACTGTCCTTCCTGTGTCTGAGGGAAGGCATGTAACTCTTGCTTATCTTCACCTGTGCTCT
AGATCCTGACCTTCTGGCAACCTCAGGGACCTTGCACCATTCTTCTCTCGCCTAAT
GGCGAGACTCAGTCTCCCTTTCCCTTTCCACTCTCCCTTTGCCATTCTTAGTATCTTTC
TACAAGCAGGTCTTCCAAAGTACTGCTTGAGGTCTGAGGGGAACATGCCTCTACC
CTACTAAAAAAGAGAAATTCCTCTGCAGAAGACCCAAGCTGACAAATCCCTTTACTG
[C, T]

AACTGCAGCTCTAGCTCCCACCATTTTCCTGTACTTACTCTCCTGCTCAGGTTCCCTGGC
ATTGCTGATGTCTTTCAGCCTTTGTGCCCTGGCCCCTTTCCTCCTCTCCCCCTCATCTAGC
ACTACCTGTCAAAATCAGGGACTTACTTTAAAATTTATCCCAAATTATCATTGCCATCAT
CTCCACTGTCACCTTATCATATGTTTGAATAGCGTTTCCATTTCCCAAATGTTTTCGCAT
GCACTTTCTCAATTGAGCCTTACGAATCCTAGAGCTGAGAAGGGTAACAATTTATGAGTC

ACTTTAGATTGAGAAGGGCTCTGACAAAGCAACATTTAAGGTGCAACCTGAGAGAATAGA AGTTAAACAGGCAGATATTGGTGAAAGAGCAGTCTAGGCAGGGAACATCATTTGCAAA GGCCCAGGGTAAAGAAGATCCTGGTAAGGAAATGACAGTGGAAGAAGGTTAGTGTAGCAG GACTGTGGCTAGGGCGGAGGCCAGGGAAGTAGTTTAGAATTTCAATGCAATAGGAAATA TGGAAGATTGAAGGCAGTTTTGCATTTTAAAATAATAATAGTTGCTATTTTAAAGCTACTT

62641 CCAGGCTGTACCAGGCACTCAGATATGACAGTGAATAGGCAACATCTTTGCCATT
GGAGAGCCTACACTGAAGTGGACATGAGGGAGTTGAAAGCAACTCTTATAGGAAATCATG
GTAAAGACGTCCAAGAGAAAGATGAAGGGCAAACACATGCACGGATGCCAAACATCT
ATCAGAGAGAAAGGAATTTTCAGACCTGACTGAATGATAGAAGGAGTTTTTGGAAAGG
AAAATAGAAGGGAAGGACAAGGGAAATTATCTGGGCAGCAATATTTATCTGCTGTGGTGC
[T,C]

63023 ATTAAAATGTACAGAAAGGAAAGCTTTGGTTCTGAGTTTGCAGGCTTCCCTGTCTTTCA
TTCCTATTGTAGAAAGCAGCTTATATAAAAAGATGTGCTGTTGGCCCTTTGAGCTGCTG
TGATTGTGTTAGGACCCCACTGGATGGTATTCGCATGAATTAATCTACTGTAGCATCTCT
ACAAATCAAGAGGCTGGCTTCTGTTTGAAATGTCCCAAGGCTTTGTGCACAGGGCAAGCT
AAATGTCTCCCTACAGTGAGACTGAAAATGCCTTGGGTGCCCTTGTCGATAGGATCTGAT
[A, G]

FIGURE 3, page 42 of 57

TGTGGTTATTTGGTTTTATGTAGAGGAGATAGAAACCAATCAGTCTAAATCATATTCTGT

GATAAGTAGGGATATTTCATAAATAATAGACGAATTGATTCATCAAGAATATACAACAAT CATAAATGTGTATGTGTCTAATAACAGAGTCTCAAATTATAATAGAAACAAAACTGACAGAA CTAAAGAGAAAAGCCAATCCTTATCTTTATCAGGTGATTTATCTTGGTG AACATTCCTTGGTGCTCTTGAAAAGAAAGTGTATTCTTGTAGTCATTGGGTATAAAATTCTA TATATGACAATGAGGTGATTGATAAAAATTCTA

66694 TCCGTTATTATCTGCAATGTCCCTATTTATCTCTGGTCATATCTTTATCTTGAAGTC
TTTTTAACTGATATGAATGTAGCCACTTCATCCTTTTTATGCTTACCATTTGCATAGTTT
ATATTTTCCATTATCTTATATTCACACTATTTATCCCTTTATACTTAAGTCCATGTCTT
GTAGACAGTATGCAGTTAATTGTGTCTTGATTATTTTACTCCTTTCTGACAATTTCTGC
CTTTCCATATAATATGCTTATCAATACAGTTGGAGTTAAATCTACCGTCTTGTTATTTGT
[C.G]

66755 TTTTAACTGATATGAATGTAGCCACTTCATCCTTTTTATGCTTACCATTTGCATAGTTTA
TATTTTCCATTATCTTATATTCACACTATTTATCCCTTTATACTTAAGTCCATGTCTTG
TAGACAGTATGCAGTTAATTGTGTCTTGATTATTTTACTCCTTTCTGACAATTTCTGCC
TTTCCATATAATATGCTTATCAATACAGTTGGAGTTAAATCTACCGTCTTGTTATTTGTC
ACATCTCCCATCTTTTGTTGTTGTTCCTCATTTCCTTGTTTATTACCTTCTTTTCAGTTA
[T.A]

FIGURE 3, page 43 of 57

ACTTCTACCACCAGGTTTTTCACACGGTTCTTCTTTCCCCATTAACAATGATCCACCATT CTCTTTCTTTATCCACTGTTACTCATCCTCATAACTGAAACATCATTTCCTAAGGATGGC [C,T]

69280 TATTGCACCACCTTGTCCCTCATCCACCTTTTTTTAGTCTTCTCTCTTTTTTAAACTT
CTACCACCAGGTTTTTTCACACGTTCTTCTTTCTCCCCATTAACAATGATCCACCATTCTCT
TTCTTTATCCACTGTTACTCATCCTCATAACTGAAACATCATTTCCTAAGGATGGCCATT
CCTGGTTCAGTCAGTCTATATTTCATCCCCCATCACATACTCTTGTTTTACCCTATATTT
TTCCTTCAAAGCACTTATTTAAGTTGTAATTATGTCTTGTTTATTTTATGTCTGCCC

70647 TCCAGTCATTTATAAAAGATGAAGAGGAGAACAAGGTAGGCCAAAGTGGCTTTGTACTAT
TAAAGGCTGCTTGATTTCTAAGTACATGTTCTTTGCCACCTTTCTGCCATTCCT
AGAAGCCATGGGTAAGTCAGCACAGGGATCTTAACATGATAACATTGGTTTTAGGAGGTC
TCGTGCATAATGGACCAGACTTAGAGCACAATGCTGTAAGGTAGTTTTAGGTGAGCAG
CAGATTCTGGCTTTAGGAGTTTATTATCAGATGCTTTTTAAACGACTTGTGGCCCAGGAT
[C.T]

CCTGCACCCATGGGAAGCATTGTAGCCTTAGAACTCTGGGAATTCTGAATATAATTCCTG
AATCAATCGTAAGGATGCATATCTGATGCTTAGTGCAAACCAAGAGGCAGAATATTTGCA
GGCAGTGTATCCTTGAAAAACAAATCTAGGTCATTTTCCTGCCATGCTTCAAGCTTACTT
TTCCATCCTTCCTGATGGTAGTACTAACTACATTTGTAGACCATTTACGTGGTCAACACT
GTGCTAAGCTGTTAGCTTCATTCTCTATGAGACAGGCACTCTTAGCCCAACTTTACAATT

71867 TCTGTCTGGCTTTTCTCAACCTTTCTCTGCACTTTCTTGGATATAATCAAAGCACTA
CCAGGAACTCCAGAGTCGGCACCTTTTCATTTTTGTGTTTTCATTTAATTATTTCTCAGC
TGCTAAGTGTTTGACTGTTTAAGGGACTCTAGTGGTAAATATTTGTCTTTAGCCTGGCAG
AAGCTGTGGTTTCCTTTGATGAGCTCACAGGTGTGGCTTTTAAGATGCTGCTGACCAGG
ACAGCTGACTGCCCCAGTGGGTCCCCAGCAGTGGCCTGCCCTTCCAGAAAG
[C, T]

71900 ACTTTCTTGGATATAATCAAAGCACTACCAGGAACTCCAGAGTCGGCACCTTTTCATTTT
TGTGTTTTCATTTAATTATTTCTCAGCTGCTAAGTGTTTGACTGTTTAAGGGACTCTAGT
GGTAAATATTTGTCTTTAGCCTGGCAGAAGCTGTGGTTTCCTTTGATGAGCTCACACGGT
GTGGCTTTTAAGATGCTGCTGACCAGGACAGCTGACTGCCCAGTGGGTGCAGTCCCCA
GCAGTGGGCTGGACCCCTTCCAGAAAGCGCTGCTGGGCCAAGAGGCTTCCTCCAACTTCC
[C, T]

GCTGCCCCCATCTAACCAACACCTCAGTCTCTTCTCCACCTGCTTCCCTGCCCTCTTCCT TTCCCTCGCAGACACTTTCTTCTGCCTGGCAAAAGGAATCTTGTTTCCATGGAAGCCTCA TTAAATCTGCATCTTGCTCAGTTTGGGTTTGATCACGGCTGCCAGAAGTATTTTTAAGCCC ATGCAGTTGCGTAATGAGATAGAGATTGGGGAAAGGGGAGGTGACTGTATAGGCAGAGG GTTTTTTTAAAAAAAAGTGAGAAAGGAAAAGGAAAACCTCTAAAGAAAAAGAGTTTTATGGA

71901 CTTTCTTGGATATAATCAAAGCACTACCAGGAACTCCAGAGTCGGCACCTTTTCATTTTT
GTGTTTTCATTTAATTATTTCTCAGCTGCTAAGTGTTTGACTGTTTAAGGGACTCTAGTG
GTAAATATTTGTCTTTAGCCTGGCAGAAGCTGTGGTTTCCTTTGATGAGCTCACACGGTG
TGGCTTTTAAGATGCTGCTGACCAGGACAGCTGACTGCCCAGTGGGTGCAGTCCCCAG
CAGTGGCTGGACCCCTTCCAGAAAGCGCTGCTGGGCCAAGAGGCTTCCTCCAACTTCCC
[G, A]

> ACTTATGCTAGGGAGTGTGATTGATGTTGCTGCTTACAGATTTCCCCTCCCACAGACCTG ATGGGGCAGCCAGGATAGTGGCAGAGAAGAAGACAGAGCAATAGCAGGAAAGAGAGACA

> > FIGURE 3, page 44 of 57

ACACTAACACATTGGAGGTTTATGTTCAAAGACGGGATCTAGGGGGTCAGAGAAAGCACA CCTACCATGTAATTGGTGCTGGAATCTGATGCCAAGTGCACCCTTGGCTTCTGAGGTTCT GAGAACTCTTGCTTGTGCTTTTCAGCCAGACTATGCCCTCACCTGCCCCTGTACTTTAAA

72992 TGTTTGCATTGGATTGTTGGAGTGTGTGTGTGTTGTTGTTTTTGTATTACAAGACA
AAGAGATTAAAAAAAACCACATGCAGCTGTCACAGCTAATGTTTATTGAACTTTTACTA
TGCCACATGGTGTTTTAAGCATTCTATATGTGTTAACTCATTTTCCCTAATTCTATGGAC
TAGACACTTAAACAGTCTCCATTGTACAAACAAGGAAACTGAGGCACAGAGAGGTTGGGA
AACTCATTTGAGGTCCTCCAGCTAATTAATAGTGGAGCCAGGTTTTGTACCCAGACAACC
[T, G]

TTTTTTTTTTTAAAGGAAAATGCTTTTCTGAGGGTGGTATCTAAATTCATAAAAATC
TTTACGATCAAGATTTTCACAAATTTCATTCTGACTCTGTTGCATTGCCCTTCTTCCCAT
ATTCCCAGTTAGTTTGTATTGATTGCTGCATCTCCCTTGAGCCCATGGTCCCCCACAACA
TTTCTTGCAGAACTGTGTCCTGCCTTCACACTGTCAGGCAGCAGGAGCCTCTCTAGCGGC
CAGCCCACAGTCCTGCAGCTCCTCCTCAGGACGTTTAATTTCCCACATTTCTATGCAGT

TTTTAAAGGAAAATGCTTTTCTGAGGGTGGTATCTAAATTCATAAAAATCTTTACGATCA
AGATTTTCACAAATTTCATTCTGACTCTGTTGCATTGCCCTTCTTCCCATATTCCCAGTT
AGTTTGTATTGATTGCTGCATCTCCCTTGAGCCCATGGTCCCCCACAACATTTCTTGCAG
AACTGTGTCCTGCCTTCACACTGTCAGGCAGCAGGAGCCTCTCTAGCGGCCAGCCCACAG
TCCTGCAGCTCCTCCTCAGGACGTTTAATTTCCCACATTTCTATGCAGTTACCTCACAG

74149 TTTGCTCAAGGTCACATAACTAGTAAGTGGGTGGAGCTGTGATGTGAAACTGGGCAGTCT
GATTCTGGGACCTGTGCTCTTAATCACCAATCTATATTGCCTCCTACTTGAAAACATCCA
GGGAAAATGTTGAGATAGATCAGCTGAAATCTTCTTGCACAGTAAAGCAGGGGCCACCTG
TCCTGGAGTTACATCATCTTGTTCATTGTCAACGATTTGTGTCAGGAACAACCAGTGACA
AGCCCAAGAACTTACCTGGGTGCTGTGACAATTGGACATGACTAGGAACAACCAGTGACA
[T. A]

TGTAGCCCATCCAAACACAGGGTAGGAAGTGGATGCTTGTCACTCTTTTTGGTTATAAG AAGCAGGAACCCAGTAAAGGCACCTTTTATATATCTATAAAGTTGAATATATAAGATATA TGGGGGCCAGGCACAGTGGCTCACACCTGTAATCCGAACATTTTTGGGAGCCCAAAGCAGG TGGATCACCTGAGGTCAGGAGTTCAAGACCAGCCTGACCAACATGGTGAAACCCCATCTT TACTAAAAATACAAAAATTAGCTGGGCGTGGTGGCACACCCTGTAGTCCCAGCTACTTG

> TAGGAAGTGGATGCTTGTCACTCTCTTTTGGTTATAAGAAGCAGGAACCCAGTAAAGGCA CCTTTTATATATCTATAAAGTTGAATATATAAGATATATGGGGGCCAGGCACAGTGGCTC ACACCTGTAATCCGAACATTTTGGGAGCCCAAAGCAGGTGGATCACCTGAGGTCAGGAGT TCAAGACCAGCCTGACCAACATGGTGAAACCCCATCTTTACTAAAAATACAAAAATTAGC TGGGCGTGGTGGCACACACCTGTAGTCCCAGCTACTTGGGAGGCTGAGGCAGGATACTTG

74918 TAACAGGTGCTGAAAACAGGAACTGGGAAGTTGCCAGTACCTTCCTGTCTTTTCCCCTGG
AACCAAACGGTTCTTACTTGCTTCTCTGCACCTCTGTCTCATTTCCCTCTCTCA
GATGATTTTCATTGCTTCACACACACATAGAAAAATCAGGATCCACCCTCCCAAGTTT
ACATATCGTTGTTTCAGGCAGCCATAGTATCCTTAAAACTCCACATTCCAGGGAGAAAGC
TTGGGTCAAGGATTCAGCCAAAGGGCAGCGAAATGGAGTAAAGATGCAACTGCCAGGTCT
[A, G]

TGGGCAGCAAGGAGGCCGGGAAGGAAGCCGCTGTTGTGGTCCAAGTGACAATTCAACAGC
TCAAAGCATAAGTAAGTTGTGTGCTTTTCACAGATGGAGAAACTGAGGCACAGAAGGAAC
CTGGCTGGGGTCCAGGTCTCTGGCCTTTGTGTCAATGCTAGGTCACTGGATGTGGCGTCT
GATTTCTACAGGAAATGTGGTTTCTCTACTTTGTCCCAGAGCCCACTCAGAGCACTGGCT
GGCCAGGGGGTCCTAGGGCCCTCTTAGGATAGTCTCAGGCCAACAGCCCCAGGACAGAAG

75386 GGATGTGGCGTCTGATTTCTACAGGAAATGTGGTTTCTCTACTTTGTCCCAGAGCCCACT

FIGURE 3, page 45 of 57

TTACTACCAGCAGCACTCAGACCCACATCTTCAGTTTAAATGTTGGAAATGGACTGTCAG
AGAACATTTAGGCCATTCATTCTGTGGGAGAGATAGGCTATGTAAAAAGATAGCCACTCC
CATGTGAACAATGTGGTTAGGATTAGAGGCATGAATATACCCCAAACCAGGGGTGTGGGA
AGGAGGTTGACACTCTAGGTGATAATACCCAGACCTTAAGGAGCTTTCTGTCTAGAGGGA
GGTATGGACATGGACAAGTAATCAACAGCTACAAAGCAGAGCTGCCAGCTCTGCAACACA

78264 ACCTTAAGGAGCTTTCTGTCTAGAGGAGGTATGGACATGGACAAGTAATCAACAGCTAC
AAAGCAGAGCTGCCAGCTCTGCAACACAAGAGCCCTGAGAGGCATGACAGGGGCAGGGTG
GGGATCCATGTGGGTCTGGATTGAAGTGAGGGGGCATCAGGAAAGCATTCCAGGAGAG
CTGAGGGACACTTGAGCACACCCTCAAAGAATGACTGGGGTCATGAGGTATACAAGGGA
GGAAGTGCACCCGAGACAGAACAATCACATAAGCAAAAATGCAGAAGAATATGAGGATC
[G,T]

80986 GCATCATATTGCATGAAAACAGCAAACGGAAGTCACAATGGCTCGACGGTGTAATGAAGC
CACACAATATGTATTAAACACATCATCTACACAGATGGATTCAAAGATACCTTCTTTGTG
TCTAAGTCCCAAATCTGTGTTTCCTGGCTCTGTTCCCTCATATCTAGTCATTCTCCAAGT
CAGCATGCCCAACTTGAAAGTGTCATTTTCAAAACCTGCTTCTTCTCTTCTGGAAGTTCT
TCCTCTGCCCATTGCTCCACAATCCCCACCTCTTTCACCCAGTAGCAAACCTTAAATTTA

83609 TTTGGGCAATTGTAGCAATTTTAAAACTATGTTAGATGGCTAGAGATTCTTGAGAATATT
TCTTTTCTTGGAAAATCATAAGGCTTTGGATAGTGGTACCTATAGAAGCTGACATCAGCA
GCAGCCTGCCTCCAGTCGATCAGGGCCTTTGGAACTCACGGGGCTCCTCTACTGACAGC
CCCATCGGTTTCCCTCCAGCACACGTAACTCAGCATTGACTCTGGGTAGTAGAGGGTGGT
TTATGGAATCTGATTCATCTCAGAAAGAGGTGGATGCAAACACATTCCCAGAGCAGAAGG
[C. T]

FIGURE 3, page 46 of 57

> GGARGA CARACACCITTOCALACTGAGTCATGGGGACAGTTTCTTCAGGAAATGGGAT CTCAGCTCTOLTT ITATOLACTACTTACATAATAAATGTTTCATTGTTTGTTTGTT ATTGTTGATTTAATAACATTCTTTTTTTTAAGAAGATTTGTAAAAACAACTGAACAAATGC AATCTCCTCCALACCIACCACACAACAAAGAAGAGAATTAGGAATAAACCCCCTTTGAGACCG TTCCTTCACCTACCTACCTACTCACTAAATACCTAAAAGCTTCAGCTAAGTAGGGTCACCCCCC

88238 CTTGTATGCAGGCTTACATAATAATATGTTTCATTGTTGTTGTTGTTGTTGATTT
AATAAGATTTTGTTACATAATAATATGTTTCATTGTTGTTGTTGTTGTTGTTGATTT
AATAAGATTTTGTTAAAAAATTTGTAAAAACAGTGAACAAATGCAATCTCCTGCC
AGAGCAGGGAGCAATAAAAAAATTAAGAATATAAGCCCCTTGGAGACGTTCCTTCACCT
ACCTGGTGGTGATATACUTAAAAGGTTCAGCTAAGTAGGGTCACCCCCCCAAGAAATTAT
TTTAAAAAAATTGAAATCTGATATTTTTAGAAAATCTTATCAAGGATATTTAATTGGACT
[A,C]

> GTACTTCAACCACTACCCTTATAGAAGTGCTGCCTAGGACCCTCTCTTCTGGCAGGTGAA GTGGAAGGAGGTTTTGCCCAAGGGAGATTCTCCCACTTCAACTTGAGTGTCTTGGT TCCGCTTTGTTTGGTTCTATTTCACCAAAGGCTTTCATCTACACATAAATTTTCTTCAGC TTTAAATAATTAGTTTTGGTAACCATTGGTATACTGGAAAGAACATTAGATTTGGAGTCC AGGTGGCTTGAGTTCAATTCTCTGCCTCTGCCATTTACCAGCTGTGTGACATTGGGCAAGT

> > FIGURE 3, page 47 of 57

92404	TGCTCTGCTACCTGCCAGCTGTTTCCCAGGGATGTGGTAAAGATGAATGGGCAAGA TCTGGGAAAGTGTTTTGAAATCCTTGATTAAAGGCCCTCCAGGCAGATGTAGAATTTTAA ATGTGTTATATTACTGCCACTATTGTTATGCTTTCTTTTATCACCCCAGAATTTCACCAT CTCCTGTTTCAGGTGAACGAGTCTGCCTGACTCTTACCTGCCTG
92672	CAAACATGTTTTTAACCAAGGGATCAGGAGGCCTTCCTGGCTGG
92684	TAACCAAGGGATCAGGAGGCCTTCCTGGCTGGCTCCTGTCAGCTGGTCATCACCTCTTA TAACTCTAGGCTTTCCCAAGCTTATTTTATT
93132	CTGCCTACTGGACAGCCACGGAGAACTTCTCCTTATCCAAGGTCGAGGAGCCCTCCGGAG TACATACTGATACCATTGGTTCTCCCACACATACCCCCATGGAGATAAAAACAGGACCCT GGAAGCCCTGTCCGTGTTTAACCAATGGGATTCAAACATGGAAATGAACTGCCCCACAAT CCACCCTGTGAGAGACCAAAGAGCAGTGTTGGATTAACAGGGAATGTTACCCTGAAAAGG CATTCAGCTTCCACTGGGGCAGCAGGTACAGTGCCAAAGATGATCCCACTTAAATTCCTAA [G, C] ACAGGAAATAAGGAAAGATGTTGTGGAAACTCAAGACCTCTCAAAGCATACTCCTTTGTA GTTCTTCCGCAGACCAGAC
93537	TGGTTCCAAACCAGCACCTGCCAAACTTCTCACCCTCTTCTGACCCTGTCCTGGGAGTTA AGAAAAAAAAATCACTTTATTGGTTGCTCCAGTTATAACTTAAACCAGACCATCAT CAAATTAAGTGACATGTACGACTGCTTATTGTATGCCAGTTACTGTGCTGTGGGGTTTTG GTTCCATTATCTCATTTAATCCTCTCAAAAACCCTGTTAGGTAGG
93557	CCAAACTTCTCACCCTCTTCTGACCCTGTCCTGGGAGTTAAGAAAAAAAA
95067	AGAGAAATGGAAGCAGGGAGATAAATTAGGTGGTTATTGCAAGAGGCCAGGTAAGAAGAG AAAGTGGTTTAAGTAGGTGGTGTGGCAGAGAAGACGGTTCCAAGCAGAGGGGGACCACG CTGACAAATAAGCGCGGGCCACTCACGCAAGCCCAACAAGGCAGAAGGCAGAAGGCAAAA GTGAAGGCCAGAGAAAACTGGACACCCTTTCCAGAGCACAGTTCAAAGGCAATGTCCT

FIGURE 3, page 48 of 57

CAAAGAAGACACTCCACCCTCCCATTTCCTCCCTATTGCCTAAAAATAAGAAGGATA
[C, T]

> CACATTAGCACAAAGGATCCACTATTCCTGCAGCCGAGCTGGGACAAGCACTTAGGCCCA CTGACTCCAACCCTTCAATAGCCTGGGACCTACGTTGTCTCCAGGTGGTATAAAACAAGA ATTTCCCCTTTGACTGGGAGAAAAAGGGAAGAACTCTAAATTGGAAAACAGGTCATCTCG AATTCTCACAGGTGGAAATTTCTGACAACCCCTTTGGGACCCACAATTCAACACACCCCA AATGGGGACAGTAGCTAACATGCAACCTGTAGGCTGTTCTGTCATCCAGTGCCACTGTGC

ATTCTTCTCACTGCCTTTCCAAGAAGGGGATTTATCAACTTCAGGGCACAGCAATCATTT ATTCCCAGACTACTGCATGCATATATATATATATTTACTTCTCTTCACTTAGAAAAAAG AGAGAATTGGAGTTGTGAATATTCCTGTCTCCCCCACCCCAGCCCCCTTGAAGTGAGTCA GGACAAACTTGGGGCCCAAATGGAGCTGTAAGTAACTGAGTCACATGCAGAGATGAAACC TTCACAGACCCACTGATATGGAGGTTGAAGATTAAATTCCCCTTTGAGAATAACTGGGTA

97271 ATTTACTTCTTGACTTAGAAAAAAGAGAGAATTGGAGTTGTGAATATTCCTGTCTCCC
TCACCCCAGCCCCCTTGAAGTGAGTCAGGACAAACTTGGGGCCCAAATGGAGCTGTAAGT
AACTGAGTCACATGCAGAGATGAAACCTTCACAGACCCACTGATATGGAGGTTGAAGATT
AAATTCCCCTTTGAGAATAACTGGGTAACACTCATACAGAGACTACTTTCAAGAAGGCCA
GATCCTCCCTCTAATGTATAGTGCAACGTTCCTAACCCTCAGCCCACTCCGTCATACCCC
[A, C]

TTTGTATGGAAAAATTTGAAAATATCAGGTGGCAGGCCAGGCATGGTAGCTCATGCCTGT
AATCCCAGCACTTTGGGAGGCCAAAGCAGGCGGATCACCTGAGGTCACGAGTTTGAGACT
AGCCGGGCCAACATGGCAAAACCCCATCTCGACTAAAAATACAAAAATTAGCTGGGTTTA
GTGGCGCATGCCTGTAATCCCAGCTACTCGGGAGGCAGGAGAATCATTTGAGCC
TGGGAGGCAAAGGTTGCAGTGAGTCAGATCATGCTACACTTCAGCCTGGGTGAGAG

97518 CCTCTAATGTATAGTGCAACGTTCCTAACCCTCAGCCCACTCCGTCATACCCCCACTCAC
ATGAATACACACATAAGCAGTAATATAAAGCACTTCCCACCATAGGGCAGCAAAGAAGGA
GGGAAATCTTTATTATGGAAGAGTGGAAGGAAGGAAGGGAAGGGAAGGGAAGGAAAAATTCTCAGGGTGAGCAGAGAATAACAGGTGGCAGGAAGAAAAATTTGAAAATATCAGGTGGCAGGCCAGGCATGGT
[G, A]

> > FIGURE 3, page 49 of 57

TCTGTTGCTGTGTAGCAAATTGTCAGAAACGTAGAGGCTTAAAGCAATACCCATTTATTA TCTCGCAAGTTCTGTATCTCAGAAGTCCAGGCAGGCTTGACTGGGTTCTCTGTCCAAGTT

AACATTTTASAAACTCTGSCTCCCCACTCACCCATAATCCTTTTAAAAACCAAATCTTGA
AGCCTTTTTTTCCCAAASGCCTTTTTTGAATAAGCACATTTATACCTAACTCATCAGACA
CCCACTTTGAAAAASACTTASCATGTGGCAAAATAGGCTGTAAATCAATCAGAACTATTC
TTTCCCACCACAAATCTTTCTCAAACACATTGGGAGAATCTGACACTGTCAGTGGTATACC
AGAGCAGACTUCTACCATTCACAAAGAGCTGACTGTTAAATGTTTAGTAATTGTGGACAT

101045 GGAAATATCASTGTTO/GTTGCTGACAGGTGGCGGTGGGGGGTTCAGTCCACGTTCA
AAGAGCCAGAAACCTGGCAGGGAAGAGATGGGGCAGTGACACCCAACCGGAAAATAAA
GGAAACTACAAGAAGAACCCAGCTAAGAGATGTGAGGGTTCTGAAAGCTCCCATGGAAAG
GTTCGCAGCTCCTCCACCTGCTCAGCTGCCCCAGGTCAAGGAAGCTCTGTGAGTG
TTAGCTGACCCGGACCAGCAAGGATACATTCAGAAGTGATGAAAGGGAACGCTTCTTGAC
[A, C]

101232 GCTCCTCCACCTCGCTCCAGCTGCCCCAGGTCAAGGAAGCTCTGTGAGTGTTAGCTG
ACCCGGAGCAGCAAGGATACATTCAGAAGTGATGAAAGGGAACGCTTCTTGACAGGGTAA
AGAGTCATTCAGTAGGAATGAGACAGGAAGAGGTCACAGAGTCAGAAGCCCAGCCTGTAC
TCAGAGATTATTTCTGGCATGGGAGGGCCGAAGGGTTAGGAGGCCACCTACTCACAATAC
AATACAGAGGCAGATCCACTTATTACCTGCCTGTGCTGCTGGGATTTCAGTGTGGAAATT
[C, G]

TGTGCCTCCTCACTGTGCTGCAGCTTGGGAATGACATCCAGAGCTTACCCACCTGCATA AGAAATAAGCTATAGGTGTAATAGGGGGACATAGGCTAAAATCCTAGCTCAGCTGCTTAA TAGCTGTGCGACTGAGCAAGTTACTTAACCTCTTTGAGCATCTGTTTTCTCATCTTTAAA ATGGAAGTAATCATAATTGACCAGGCCCAGTGGCTCACACCTATAATCCCAGCACCTTGG.AAGGCCCAGGGCCCAGTGGTTAATCCCAGCACCTCT

> ACATCCAGAGCTTACCCACCTGCATAAGAAATAAGCTATAGGTGTAATAGGGGGACATAG GCTAAAATCCTAGCTCAGCTGCTTAATAGCTGTGCGACTGAGCAAGTTACTTAACCTCTT TGAGCATCTGTTTTCTCATCTTTAAAATGGAAGTAATCATAATTGACCAGGCCCAGTGGC TCACACCTATAATCCCAGCACCTTGGAAGGCCGAGGCCAGTGGATTGCTTGAGCCCAAGA GTTTGAGACCAGCATGGTGACACCTCGTCTCTAGAAAAAATACAAAAATTAGCCAGGCAT

101290 TGACCCGGAGCAAGGATACATTCAGAAGTGATAAGGGAACGCTTCTTGACAGGGT AAAGAGTCATTCAGTAGGAATGAGACAGGAAGAGGTCAGAAGCCCAGCCTGT

FIGURE 3, page 50 of 57

ACTCAGAGATTATTCTGGCATGGGAGGGCCGAAGGGTTAGGAGGCCACCTACTCACAAT ACAATACAGAGGCAGATCCACTTATTACCTGCCTGTGCTGCTGGGATTTCAGTGTGGAAA TTCTGTGCCTCCTCACTGTGCTGCAGCTTGGGAATGACATCCAGAGCTTACCCACCTGC

TAAGAAATAAGCTATAGGTGTAATAGGGGGACATAGGCTAAAATCCTAGCTCAGCTGCTT
AATAGCTGTGCGACTGAGCAAGTTACTTAACCTCTTTGAGCATCTGTTTTCTCATCTTTA
AAATGGAAGTAATCATAATTGACCAGGCCCAGTGGCTCACACCTATAATCCCAGCACCTT
GGAAGGCCGAGGCCAGTGGATTGCTTGAGCCCAAGAGTTTGAGACCAGCATGGTGACACC
TCGTCTCTAGAAAAAATACAAAAATTAGCCAGGCATGGTGGCAGGTGCCTGTAGTCTTAG

> GCTAAAATCCTAGCTCAGCTGCTTAATAGCTGTGCGACTGAGCAAGTTACTTAACCTCTT TGAGCATCTGTTTTCTCATCTTTAAAATGGAAGTAATCATAATTGACCAGGCCCAGTGGC TCACACCTATAATCCCAGCACCTTGGAAGGCCGAGGCCAGTGGATTGCTTGAGCCCAAGA GTTTGAGACCAGCATGGTGACACCTCGTCTCTAGAAAAAATACAAAAATTAGCCAGGCAT GGTGGCAGGTGCCTGTAGTCTTAGCTACTCGGTAGGCTGAGGTGGGAAGATTATATGAGC

> CCAGACATGTATTTCCTAATCGTCTCCAGGTTGTTTGATAGAAGATCTCCTGGGAGCAGG TTTCCGCAGCAGCTCAGCCAGGTCTGTTCTGGGAACGCTGTGTGCATTGGCACCTCCCTT GGCAGAAAGCTTGGAGGAAAGGCAGGTGCAGGTCCTGGAGCCTCTGACAGCATTACTGGC TCTAGGAGTAGCTCCTCAGGATAATCTGTCCCCATGACCATTAAGTAACTGCCACTGTGC GGGAAGAACACTGGAAATGGGGGGCCCAAAAAAATCTGAAAACCCTCACTTGAACCAGT

> TAGATTTTGAGAGAAATCTCTCTGTTACCACCCCTTAACATTCCAACCCCCTCTAATAGCC CATTTAGGATTTATCATACTGTTTCATCCAAACCTTTCATGACCTGATTTCTATTTCCAG CTTCAACCACCCCTTGGGTCACCACCTGTACTTATTGAGTTTCCCTAGTTTTCTGAATTA ATGACTGAAGATGATAAGCTTCCCTTACATATGACTCTCAAACCACCAAACTGGGATTGT TGTTACTCTTAGTGATAATGGTTGCTATTTATGAAACTTTTAATAGGGAACACAAACCCT

105266 AGGCCAGAGCATCATGGCCTTTCACAAGTTGAAGAGCCACGGGCTTTCTACGGTAGCCAG
CCACGCTTTTCCATGACTGGGGTGGGTGGCAAGTGATGAGGGGTTTGGAGTTCATGTGG
TGGGTGGCAGGGACCAGGTGTCTTGGTAACTGCTGTTGCATTCACTTCAGGAGCAAAGG
ACCAGATCTGATTCTGCAGGATCAACAATATGGACACTGCAGGCTCTGTAGACATCCAAA
GCTCTAATGGTGACTTGGGGAAGCTCAGGAGGGCAGGGAGGTTGTACCCATTTAGAATGT
[A, G]

105338 ATGACTGGGTGGCTAGCCAAGTGATGAGGGTTTGGAGTTCATGTGGTGGGGTGGCAGG
GACCAGGTGTCTTGGTAACTGCTGTTGCATTCACTTCAGGAGCAAAGGACCAGATCTGAT
TCTGCAGGATCAACAATATGGACACTGCAGGCTCTGTAGACATCCAAAGCTCTAATGGTG
ACTTGGGGAAGCTCAGGAGGCAGGAGGTTGTACCCATTTAGAATGTAAAGATTCCTAT
TTTATAAAAAAAGAAAAAAAGGAAGACTGAAGGCCTCAGTCTCCTCCAACAAAGCCAGGCTG
[T.C]

**AATAAATAAATAAATGAAGGAAGAAAAAAAGAAGAAGAAATGCAGAACAGGGTGACTAAA** 

FIGURE 3, page 51 of 57

ATTGGCATGTATTTTAAATGTTTATATTAACAAACTAACACCTTTTAACATGAAAAGCA ATATAATTGTGCTAGCCACAAAATCATCGTAGGACTGAGAAAGGAATCGTGATTCTGAGA GCCCTAGAGTTAATGTGATCCAGCTGGCTCATCCCTGTGACTGCAGAAGCCTGTTTGGAG ATAGTGTCAGTAGCTTTTCAGGCCCTCTGTGAATTGCCAGAATGTGTGACATGAGCCAAA

105928 AAAATTGGCATGTATTTTTAAATGTTTATATAACAAACTAACACCTTTTAACATGAAAA
GCAATATAATTGTGCTAGCCACAAAATCATCGTAGGACTGAGAAAGGAATCGTGATTCTG
AGAGCCCTAGAGTTAATGTGATCCAGCTGGCTCATCCCTGTGACTGCAGAAGCCTGTTTG
GAGATAGTGTCAGTAGCTTTTCAGGCCCTCTGTGAATTGCCAGAATGTGTGACATGAGCC
AAATTTCCCCCCAGCATCCCCGCCGCCGCCACCACCCCCCGACCCAACCCTCCCGCCG
IG.AI

CTCCCATAGAATAGTCACTGCCATACAGAAAAAGAGAAGTTCTACTATTTCTGGGCAAGA
TTTCCACAAACCAGTTTGTCCCTTTCTGCTTTCATGAAATAAACCATTTGGATCAACGTC
AGCTGATTGCAAAAATTTTCCCTTGTCTCAAAAGCAAGACTGATAAGGAAGCAAACATGG
GAGGACCTTAGTGGCCGAGCCTTTATGTGTATGTTATTTCATTGCTCTCATAACTGCCCT
GGGATGCTGTAAGCATGATTCATCCTGTTTTTTTTATCAGTTAAATTATGTATCCAAGATT

TCTTTTGTCCGCTGAGCAAGGTATAAAAAGATGTCAAAAGAAGTACCCAAAAAGGTAATA AAAATGTACAGTCGTGCATCACTTAGCAATAAGGATACATTCTGAGGAAGGTGTCCTTAA GCAATTTTGTCATCGTGGGAAAATTATAGAGTGTACTTTCACAAACCTAGATGGTGTAGC CTACAACACCTGGACTATGTGGGCCTATTGCTCCTAGGCTACAAACCTGTACAGCATG TGCTTGTACTGAATATTGCAGGCAACTGTAGCACATGTTTGTGTATCTAAACACAT

108062 AAGGTAATAAAAATGTACAGTCGTGCATCACTTAGCAATAAGGATACATTCTGAGGAAGG
TGTCCTTAAGCAATTTTGTCATCGTGGGAAAATTATAGAGTGTACTTTCACAAACCTAGA
TGGTGTAGCCTACAACACCTGGACTATGTGGGCCTATTGCTCCTAGGCTACAAACCTG
TACAGCATGTGCTTGTACTGAATATTGCAGGCAACTGTAGCACAATGGTATTTGTGTATC
TAAACACATCTAGACATAGAAAAGGCACAGTAAAAATATCGTAGTATATAGCCTTATGGG
[G, A]

FIGURE 3, page 52 of 57

> TCTCTCTCTCTCTCTCTCTCTCTCTGTCTGGTTTTCCTTCCTCATAAATACTTTT

111484 ACAGGCCTTCTCAGTGTGATTGGTCATTGTCTGTGGGGACTCTCCTGCAGAG
CTGACCACTTCTGTGCCTGCGCTGGTTTGGACACACCTGATGCTCTAGGGGCAGACTCC
TCTCCTTCTTCACTGCTGCTTCTTCTTCGTCACCACTCAATAAAACGTTGCCCTCAGCCTG
ACTGCCAAAAAGTGCTGGAAGAAAAATTATCTCTGGTTCTATTGTTTCCCACATTGTA
TTCTTGCCCAACTTCCAGTTCTTGCCACCAACAATATTCTCAGAGGTTGCCTCAGCACCT
[G.T]

GTTTATGTATCATGCAGACTCTGGATCCACATATATCTCAGTGGCTGTGAATATAGGATG ATTGATCACAGGCCTGAGTTGCATTCCTACAGATTCTTAGGAAAAAAATTGATTCACAGA CATGTCCCCCCTGGTTCCCCCACAACACACCTCCTTCCTCAGCAATCTCTATCAGTCAC CAACTACACGTTGAATATGTGGCAAGCTCTTCCCAGACCTTTATCTGAGAGCCAAGGAGT GAGGGGCTGTACTAAGATATCATAGAAATGAAAATGTGGTGTGTCACAAGTTTCCTTAAT

FIGURE 3, page 53 of 57

[C,G]

CAAGGAGTGAGGGGCTGTACTAAGATATCATAGAAATGAAAATGTGGTGTGTCACAAGTT TCCTTAATTCTTAGATCTTAAACTCTAAGAGGGTTCAGCATAAGTACAAATTCAAGGGCT AGAGACAACCTGTATTGGGTGTGTCTTTAACTCAGTTTCCCAATCCACATAGGGACCTTG CATTTGTCATCTCTCATCTATGTATAGCTGTTGGTATGACAGTTTCTCTGTTCCAGAATA CCTGAACTCTGACTTAGCCTGTCCTTTCTGAAACAGAAAAATCACCCAACCAGAGATCTA

114486 CCCCATGGTCATTTTGCCACTCATAAGTTAGCTACTCTGGCAGGGTTGCAACTTACACA
GTTTTCATGATAACTGGATTCTCACTCCTTTTTTTACAGAATGGATGTGATAACCTGGTA
TCCTACACAGTCATGAGTGACCAACCTACCCATTTGGTTCCCCATCCTCATTCCTCCATT
CCTAGCCCTAGGGTAGCCGGGAAAGCATAGGAGCAAATGCCCTTACCAGGGCCCTGGTGC
TCAGCAGCCTCTCCGGCTGCTCACACCTCTTGCTGCTCTCTGTGCATGCTCCAAAGGCT

CTTTTTGCGTATGCTGAGCTCTCACCTACTAAGCTCTCTGCTTTCCTTATGCTGCC
AGCAACCACAAAACCTGGTGATACTTTCAAGATGGGACATTAATGCTCTTTCCTTTCTT
TCTTCCATTTTCTGGTATCCATTTGCAAACAGCGCTCCTGTTATCTCCAGGTAAGAGGT
GTCTTGTCCCCCTCTTTCTTTCCACTTCTTGCCAGTGCCATTATTTGGTTTAAGACCAA
TGTCCTTTGATTTATTGAATAAGAACTGCAGGCTCAAGTTAACCTGACAATTTCTCCCAA

AGAGAATTTATTTTGAAGAGATTCTCATGCAGAATCTAGGTGCTATAGAGGACGTACACC
TACTTTGAGAGTATGCTTGCATGAGTGGAAACCAATCATAAACAACATTCAACTTCATGA
GCAGATATGAAAGCATTTCAGCATATCTAGCAATACTATTAGACTCTTTGTGCAAGCAGAG
TGGCCTACACAGACAGTTTCAATATATTTTTAAAAGAACGTCTTACATTTCATCAGTCCT
TTGAACACAGAAAAAAAATGTTAAGGCCACTTAAGAGGCAAAACATCTTACAGAGTTCATT

ATATTCAAAGTCACCTACAGGCTACATCTTGGGTTCAGGAAGGGGCGGTGTACATAGTAA GGACATACGCCTTCTGGGAGCCTTAAACAAAAAAAAATGTAGGTAACTCCTACATTT TTCTTTTGTGGAAAAAAACAAGTTACTCCAGCTTCCTTGGCTTTTTTGTTCTTTTTTTATA CCAACAAAATAAGGGCTATCCTCAACCCTCTGTTCTTCATTCTTCTCCCAGGGTATTGAT TTCATAACATTGGGTTTTTCTTCTCTACTTCACTCATCCTCTTGCCTGTGAAGGTATGTA

115668 GAGTATGCTTGCATGAGTGGAAACCAATCATAAACAACATTCAACTTCATGAGCAGATAT
GAAAGCATTTCAGCATATCTAGCAATACTATAACTCTTTGTGCAAGCAGAGTGGCCTAC
ACAAGACAGTTTCAATATTTTTAAAAGAACGTCTTACATTTCATCAGTCCTTTGAACAC
AAGAAAAAATGTTAAGGCCACTTAAGAGGCAAAACATCTTACAGAGTTCATTGATATTCA
AAGTCACCTACAGGCTACATCTTGGGTTCAGGAAGGGGCGGTGTACATAGTAAGGACATA
[],,C]

CTCCAGCTTCCTTGGCTTTTTGCTTCTTTTTTATACCAACAAAATAAGGGCTATCCTCAA CCCTCTGTTCTTCATCTTCTCCCAGGGTATTGATTTCATAACATTGGGTTTTTCTTCTC TACTTCACTCATCCTCTTGCCTGTGAAGGTATGTAAGGCTTCTTTGTTCCAACTCTTTCC TCCACCCGCCCCCCCCACATAAATGCATAACAAAGATTGTGATTTAATTTAAGTTTCTT

FIGURE 3, page 54 of 57

## TCTACTTTAACATATTTGCAAACATCAATAGAAGCTAAAATGGGAAAAAGGAAATGTTT

117230 AATAATACTGTCGCTGCTAAGATAGGCATTGTGATATGGTGCTTAAACCTGCAAGTAAAG
GAAAAGAGTATGGAATCTGTGTGTCTTTTTCTAAGGCCTTTTTCCCAGAGTAGCTTGCAG
TCTGGCTTCTAGGGTTGCTGGCCTATAGCCAGAACCCTAGATTCACCCAGATTTACCTTC
AGAATTAACTAATCAGAGACTCAAATTCAATAGACTAAATGAAGTCAGGCTGCTAGAGGA
TGTCTGCTGACTTGGACATATGCAGAAAGACATTGGATCCTTGAGAAAAACATTGTTTCCAA
[A,C]

121926 TTGGTCTCAAAGATTCAGTCACAGCTGTTGTTTTCGTGGCATTTGGCACCTCTGTCCCAG
GTGAGAGTGAGAGGTGCTTGAATTTGCAAAGAGGATTTTACCTGGTTCAAATGACCCCTG
GACTCCATCTCATTATCTTCCACCACCATCTCAGATCTGAACTTAACAGAGCCTCTGCCCT
TAAAGTGCACAAAAGTCAATCAAAGAGATGAATAATGACATTAGTAATGACAGCTAATAT
TTCTTGAGCACTTTCAATGTGACAGACACCATGTGTGTCAGCAATTTACAAT

ATATTGTTCTCCTTCTATTTACCTCTGGCGATCTCTGAGAGGTTAAAGATTAGCCAGCTC
AAAGATATCAAAGGAGAAATGCCCACATACATTCTTGGCCTCCTCTACTTGGAAGGACAC
TGTGAGTACAAAGTATCTCCTAGCAGGACAGCCAAAGGAAGTTCCACAGCTTTTATCTTT
TTATAGGATGAATTACATACTCTTTCTTTTTCTTAGGAACACTCAGAGACAAAAAGGAAAG
GAGCGGACATTCCTTTACTCATTGAACAAATATTTACTGAGCACCTATTATGCCTGTTAC

FIGURE 3, page 55 of 57

TTCACATCAAATTCTAGGAAAACCTCTTTCCAAAACCCCAGCGCAGGCCAGCGGTATTAT
TTGTCCATTAGTGATGCAAGAGATTTAGCTATCGTGGAAATGCATCAGAAGGTTGGAAAT
[T.C]

123366 GGTGGGAAGAAATATCAGAGGATCAGAAGCAAAAAACAACAATAACAACAGAAACAAAAA CAAACAAACAAAAAACAAAAAACAAGGCCATAGGCAAGAAAGGGTAAGAGGTTTTCTCTGGG AGATCTAAAAAAAAATGGCAATAATGAGGTAAGCCAGGCAGATACCTTTGGGCATCTCCAA GTCCTTGCAATTGGCCAAGAACAACAACATTTGAGGCTTTAAGAAGGTTACCCT GTGATCCACTCATCTGATTTAGTGGCTTTGGCTGAAGCTCTTTGGATATAGTTGAAGGTA [C, T]

GGAAAGGGTCCTTACATGAGGACTTTAGGGTCAAGTCTCTTGCTAACATCCTATGTGACC
TTGGGTAAATTCTTTGACCCTTATTTTTCTTACCTGTAAAATAAAGAATTGGGCTAGAT
GTCTCTGACAGTCCTCCCTGTATCTACAATCTGTGCCAAGATCTAAAGTCAAACACCCTG
CAAGGCCCTGTGATACATATATAAACCACAAAGACAGAGCCCCGTCTTCCTTGAGTCCAC
AGTTCACCCTGCATGTCCCCATCATGGTTCCCCAACATGTCCTCTGTCCCCAAAATCCAG

126043 AAAGCATTTTTACAAGATAGGAACTGGAATTCCTCATTTCTCCCATGTTCCTGCTTGTTC
TTAAACTTCATGAAGCTATTTTTCCAGCCTATGGGGTAGTTCTTGCTCCAGTAAGAGGAA
TCTTAGTTGTCATAATCCCTTGGAGCCTGGGTTTTTGGAGAAAGAGATCTCCGTGCCCTA
CAGACCTTTTCTCAACGAATGTGGGAAGGACCTGGCTTTAAAACACGCACACAAACACAC
AAATAAACAGACATAAGATGTCATCACGAAACTGCCCACGGATCTTTAGGCTTTCTGCAT
[T.C]

126064 AACTGGAATTCCTCATTTCTCCCATGTTCCTGCTTGTTCTTAAACTTCATGAAGCTATTT
TTCCAGCCTATGGGGTAGTTCTTGCTCCAGTAAGAGGAATCTTAGTTGTCATAATCCCTT
GGAGCCTGGGTTTTTGGAGAAAGAGATCTCCGTGCCCTACAGACCTTTTCTCAACGAATG
TGGGAAGGACCTGGCTTTAAAACACGCACACAAACACACAAATAAACAGACATAAGATGT
CATCACGAAACTGCCCACGGATCTTTAGGCTTTCTGCATTGACATAAATACATTTTCTAA
[-, G]

> TGTGGGCTTCTTGTGAGGAGACGTGACTCAGGTGAAGGTGTCACCTCCTCTCACACTCAG GTGCCAATGTGTCAGACCCAGTATATTCTAAGCAAAAATACTTCAGGAAAATGCCACTTG

> > FIGURE 3, page 56 of 57

Chromosome map: Chromosome 14

FIGURE 3, page 57 of 57

## SEQUENCE LISTING

```
<110> PE CORPORATION (NY)
<120> ISOLATED HUMAN TRANSPORTER PROTEINS,
  NUCLEIC ACID MOLECULES ENCODING HUMAN TRANSPORTER PROTEINS,
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<151> 2000-17-10
<150> 09/804,474
<151> 2001-13-03
<160> 4
<170> FastSEQ for Windows Version 4.0
<210> 1
<211> 2782
<212> DNA
<213> Human
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gtgccaagca cagggcagaa caatgagtcc tgttcagggt catcggactg caaggagggt 180
gtcatcctgc caatctggta cccggagaac ccttcccttg gggacaagat tgccagggtc 240
attgtctatt ttgtggccct gatatacatg ttccttgggg tgtccatcat tgctgaccgc 300
ttcatggcat ctattgaagt catcacctct caagagaggg aggtgacaat taagaaaccc 360
aatggagaaa ccagcacaac cactattcgg gtctggaatg aaactgtctc caacctgacc 420
cttatggccc tgggttcctc tgctcctgag atactcctct ctttaattga ggtgtgtggt 480
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atgttcatca tcattggcat ctgtgtctac gtgatcccag acggagagac tcgcaagatc 600
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Arg Phe Met Ala Ser Ile Glu Val Ile Thr Ser Gln Glu Arg Glu Val
                                 105
Thr Ile Lys Lys Pro Asn Gly Glu Thr Ser Thr Thr Ile Arg Val
                             120
Trp Asn Glu Thr Val Ser Asn Leu Thr Leu Met Ala Leu Gly Ser Ser
                         135
                                            140
Ala Pro Glu Ile Leu Leu Ser Leu Ile Glu Val Cys Gly His Gly Phe
                    150
                                        155
Ile Ala Gly Asp Leu Gly Pro Ser Thr Ile Val Gly Ser Ala Ala Phe
                165
                                    170
Asn Met Phe Ile Ile Gly Ile Cys Val Tyr Val Ile Pro Asp Gly
            180
                                185
Glu Thr Arg Lys Ile Lys His Leu Arg Val Phe Phe Val Thr Ala Ala
        195
                             200
                                                205
Trp Ser Val Phe Ala Tyr Ile Trp Leu Tyr Met Ile Leu Ala Val Phe
                        215
                                            220
Ser Pro Gly Val Val Gln Val Trp Glu Gly Leu Leu Thr Leu Phe Phe
                    230
                                        235
Phe Pro Val Cys Val Leu Leu Ala Trp Val Ala Asp Lys Arg Leu Leu
                                    250
Phe Tyr Lys Tyr Met His Lys Arg Tyr Arg Thr Asp Lys His Arg Gly
                                265
Ile Ile Ile Glu Thr Glu Gly Glu His Pro Lys Gly Ile Glu Met Asp
                            280
Gly Lys Met Met Asn Ser His Phe Leu Asp Gly Asn Leu Ile Pro Leu
                        295
Glu Gly Lys Glu Val Asp Glu Ser Arg Arg Glu Met Ile Arg Ile Leu
                    310
                                        315
Lys Asp Leu Lys Gln Lys His Pro Glu Lys Asp Leu Asp Gln Leu Val
                325
                                    330
Glu Met Ala Asn Tyr Tyr Ala Leu Ser His Gln Gln Lys Ser Arg Ala
                                345
Phe Tyr Arg Ile Gln Ala Thr Arg Met Met Thr Gly Ala Gly Asn Ile
                            360
Leu Lys Lys His Ala Ala Glu Gln Ala Lys Lys Thr Ala Ser Met Ser
                        375
                                            380
Glu Val His Thr Asp Glu Pro Glu Asp Phe Ala Ser Lys Val Phe Phe
                                        395
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Asp Pro Cys Ser Tyr Gln Cys Leu Glu Asn Cys Gly Ala Val Leu Leu 410 Thr Val Val Arg Lys Gly Gly Asp Ile Ser Lys Thr Met Tyr Val Asp 420 425 Tyr Lys Thr Glu Asp Gly Ser Ala Asn Ala Gly Ala Asp Tyr Glu Phe 440 Thr Glu Gly Thr Val Val Leu Lys Pro Gly Glu Thr Gln Lys Glu Phe 455 460 Ser Val Gly Ile Ile Asp Asp Asp Ile Phe Glu Glu Asp Glu His Phe 470 475 Phe Val Arg Leu Ser Asn Val Arg Val Glu Glu Glu Gln Leu Glu Glu 485 490 Gly Met Thr Pro Ala Ile Leu Asn Ser Leu Pro Leu Pro Arg Ala Val 500 505 510 Leu Ala Ser Pro Cys Val Ala Thr Val Thr Ile Leu Asp Asp Asp His 520 Ala Gly Ile Phe Thr Phe Glu Cys Asp Thr Ile His Val Ser Glu Ser 535 Ile Gly Val Met Glo Val Lys Val Leu Arg Thr Ser Gly Ala Arg Gly 550 555 Thr Val Ile Val ito the Arg Thr Val Glu Gly Thr Ala Lys Gly Gly 565 570 Gly Glu Asp Pho Glu As; Thr Tyr Gly Glu Leu Glu Phe Lys Asn Asp 580 585 Glu Thr Val Lys Thr The Arg Val Lys Ile Val Asp Glu Glu Glu Tyr €00 Glu Arg Gln Glu Asn the Phe Ile Ala Leu Gly Glu Pro Lys Trp Met 615 620 Glu Arg Gly Ile Ser Ala Lec Leu Leu Ser Pro Glu Val Thr Asp Arg 630 635 Lys Leu Thr Met Glu Glu Glu Ala Lys Arg Ile Ala Glu Met Gly 645 650 Lys Pro Val Leu Cly Glu His Pro Lys Leu Glu Val Ile Ile Glu Glu 660 665 Ser Tyr Glu Phe Lys Ser Thr Val Asp Lys Leu Ile Lys Lys Thr Asn 680 Leu Ala Leu Val Val Gly Thr His Ser Trp Arg Asp Gln Phe Met Glu 700 Ala Ile Thr Val Ser Ala Ala Gly Asp Glu Glu Glu Asp Glu Ser Gly 710 715 Glu Glu Arg Leu Pro Ser Cys Phe Asp Tyr Val Met His Phe Leu Thr 725 730 Val Phe Trp Lys Val Leu Phe Ala Cys Val Pro Pro Thr Glu Tyr Cys 740 745 His Gly Trp Ala Cys Phe Val Val Ser Ile Leu Ile Ile Gly Met Leu 760 Thr Ala Ile Ile Gly Asp Leu Ala Ser His Phe Gly Cys Thr Ile Gly 775 780 Leu Lys Asp Ser Val Thr Ala Val Val Phe Val Ala Phe Gly Thr Ser 790 795 Val Pro Asp Thr Phe Ala Ser Lys Ala Ala Ala Leu Gln Asp Val Tyr 810 Ala Asp Ala Ser Ile Gly Asn Val Thr Gly Ser Asn Ala Val Asn Val 825 Phe Leu Gly Ile Gly Leu Ala Trp Ser Val Ala Ala Ile Tyr Trp Ala 840 Met Gln Gly Gln Glu Phe His Val Ser Ala Gly Thr Leu Ala Phe Ser 855 Val Thr Leu Phe Thr Ile Phe Ala Phe Val Cys Leu Ser Val Leu Leu 870 Tyr Arg Arg Pro His Leu Gly Glu Leu Gly Gly Pro Arg Gly

Cys Lys Leu Ala Thr Thr Trp Leu Phe Val Ser Leu Trp Leu Leu Tyr 900  $\cdot$  905  $\cdot$  910  $\cdot$  Val Leu Phe Ala Thr Leu Glu Ala Tyr Cys Tyr Ile Lys Gly Phe 915  $\cdot$  920  $\cdot$  925